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- (54) Title: MAMMALIAN CHEMOKINE REAGENTS
- (57) Abstract

Novel chemokines from mammals, reagents related thereto including purified proteins, specific antibodies, and nucleic acids encoding said chemokines. Chemokine receptors are also provided. Methods of using said reagents and diagnostic kits are also provided.

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MAMMALIAN CHEMOKINE REAGENTS

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FIELD OF THE INVENTION

The present invention relates to compositions related to proteins which function in controlling physiology, development, and/or differentiation of mammalian cells, e.g., cells of a mammalian immune system. In particular, it provides proteins and mimetics that regulate physiology, development, differentiation, and function of various cell types, including hematopoietic cells. It also provides receptor reagents for chemokine-like proteins.

BACKGROUND OF THE INVENTION

The circulating component of the mammalian 20 circulatory system comprises various cell types, including red and white blood cells of the erythroid or the myeloid cell lineages. See, e.g., Rapaport (1987) Introduction to Hematology (2d ed.) Lippincott, Philadelphia, PA; Jandl (1987) Blood: Textbook of Hematology, Little, Brown and Co., Boston, MA.; and Paul 25 (ed.) (1993) Fundamental Immunology 3d ed., Raven Press, N.Y. Progression through various stages of differentiation is regulated by various signals provided to the cells, often mediated through a class of proteins 30 known as the cytokines. Within this group of molecules as a further group known as the chemoattractant cytokines, or chemokines. See, e.g., Schall (1994) "The Chemokines in Thomson (ed.) The Cytokine Handbook (2d ed.) Academic Press; and Schall and Bacon (1994) Current 35 Opinion in Immunology 6:865-873.

Although the full spectrum of biological activities of the chemokines has not been extensively investigated, chemoattractant effects are recognized. The best known biological functions of these molecules relate to

chemoattraction of leukocytes. However, new chemokines are being discovered, and their biological effects on the various cells responsible for immunological responses are topics of continued study.

Certain chemokine receptors have also been characterized. See, e.g., Samson, et al. (1996)

Biochemistry 35:3362-3367; and Rapport, et al. (1996) J.

Leukocyte Biology 59:18-23.

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These observations indicate that other factors exist whose functions in hematopoiesis, immune development, and 10 leukocyte trafficking were heretofore unrecognized. These factors provide for biological activities whose spectra of effects are distinct from known differentiation, activation, or other signaling factors. 15 The absence of knowledge about the structural, biological, and physiological properties of the regulatory factors which regulate hematopoietic cell physiology in vivo prevents the modification of the effects of such factors. Thus, medical conditions where regulation of the development or physiology of relevant 20 cells is required remains unmanageable.

SUMMARY OF THE INVENTION

The present invention is based, in part, upon the 25 discovery of new genes encoding chemokines, and new genes encoding various receptors for chemokines. It embraces agonists and antagonists of the chemokines. In particular, sequences of various chemokines, e.g., designated Thymus Expressed ChemoKine (TECK); MIP-3a; MIP-3 β ; and chemokine receptors designated "dendritic 30 cell receptor for chemokine" (DC CR) and "monocyte/dendritic cell receptor for chemokine" (M/DC CR); and mutations (muteins) of the respective natural sequences, fusion proteins, chemical mimetics, 35 antibodies, and other structural or functional analogs are provided. It is also directed to isolated genes encoding respective proteins of the invention. Various uses of these different protein or nucleic acid compositions are also provided.

The present invention provides a substantially pure or isolated polypeptide comprising a segment exhibiting sequence homology to a corresponding portion of a mature TECK, MIP-3 α , MIP-3 β , DC CR, or M/DC CR, wherein the homology is at least about 70% identity and the portion is at least about 25 amino acids. Preferably, the protein further comprises a second segment exhibiting at least about 90% identity over at least 9 amino acids; or at least about 80% identity over at least 17 amino acids. In other preferred embodiments, the polypeptide: is from 10 a warm blooded animal selected from the group of birds and mammals, including a mouse or human; comprises a natural sequence set forth in SEQ ID NO: 2, 4, 6, 8, 10 or 12; exhibits a post-translational modification pattern 15 distinct from a natural form of the polypeptide; is made by expression of a recombinant nucleic acid; comprises synthetic sequence; is detectably labeled; is conjugated to a solid substrate; is conjugated to another chemical moiety; is a fusion protein; is in a denatured 20 conformation, including detergent denaturation; further comprises an epitope tag; is an immunogenic polypeptide; has a defined homogeneous molecular weight; is useful as a carbon source; is an allelic variant of SEQ ID NO: 2, 4, 6, 8, 10, or 12; is a 3-fold or less substituted form of a natural sequence; is in a sterile composition; is in 25 a buffered solution or suspension; is in a regulated release device; comprises a post-translational modification; is in a cell; or is in a kit which further comprises instructions for use or disposal of reagents 30 therein.

In other aspects, the invention provides an isolated or recombinant nucleic acid encoding such protein, where the portion consists of sequence from the coding region of SEQ ID NO: 1, 3, 5, 7, 9, or 11. Other aspects include such nucleic acids which: exhibit at least about 80% identity to a natural cDNA encoding said segment; is in an expression vector; further comprises a promoter; further comprises an origin of replication; is from a natural source; is detectably labeled; comprises

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synthetic nucleotide sequence; is less than 6 kb; is from a mammal; comprises a natural full length mature coding sequence; is in a kit, which also comprises instructions for use or disposal of reagents therein; is a specific hybridization probe for a gene encoding the protein; is a PCR product; or is in a cell. The invention also provides a method of using a purified nucleic acid by expressing the nucleic acid to produce a protein.

Alternatively, the invention provides an isolated or 10 recombinant nucleic acid which encodes at least eight consecutive residues of SEQ ID NO: 2, 4, 6, 8, 10, or 12. Preferably, that nucleic acid encodes at least: twelve consecutive residues from SEQ ID NO: 2, and further comprises a coding sequence of at least 17 nucleotides 15 from SEQ ID NO: 1; twelve consecutive residues from SEQ ID NO: 4, and further comprises a coding sequence of at least 17 nucleotides from SEQ ID NO: 3; twelve consecutive residues from SEQ ID NO: 6, and further comprises a coding sequence of at least 17 nucleotides 20 from SEQ ID NO: 5; twelve consecutive residues from SEQ ID NO: 8, and further comprises a coding sequence of at least 17 nucleotides from SEQ ID NO: 7; twelve consecutive residues from SEQ ID NO: 10, and further comprises a coding sequence of at least 17 nucleotides from SEQ ID NO: 9; or twelve consecutive residues from 25 SEQ ID NO: 12, and further comprises a coding sequence of at least 17 nucleotides from SEQ ID NO: 11. preferred embodiments, the nucleic acid: exhibits at least about 80% identity to a natural cDNA encoding the 30 segment; is in an expression vector; further comprises a promoter; further comprises an origin of replication; encodes a 3-fold or less substituted sequence from a natural sequence; is from a natural source; is detectably labeled; comprises synthetic nucleotide sequence; is less than 6 kb; is from a mammal; is attached to a solid .35 substrate, including in a Southern or Northern blot; comprises a natural full length coding sequence; is in a cell; or is in a detection kit, which also comprises instructions for use or disposal of reagents therein.

Further embodiments include a nucleic acid which hybridizes under stringent wash conditions of 55° C and less than 150 mM salt to the nucleic acid; while preferred embodiments include those which exhibit at least about 85% identity over a stretch of at least about 30 nucleotides to a primate sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11; or where the identity is at least 90%; or the stretch is at least 75 nucleotides; or where the identity is at least 100 nucleotides.

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In other embodiments, the invention provides a binding compound comprising an antigen binding fragment from an antibody which binds to a mature TECK, MIP-3 α , MIP-3β, DC CR, or M/DC CR protein. In various embodiments, the binding compound is one wherein: the 15 polypeptide is a mouse or human protein; the antibody is raised against a mature peptide sequence of SEQ ID NO: 2, 4, 6, 8, 10 or 12; the antibody is a monoclonal antibody; the binding compound is attached to a solid substrate; 20 the binding compound is in a sterile composition; the binding compound binds to a denatured antigen, including a detergent denatured antigen; the binding compound is detectably labeled; the binding compound is an Fv, Fab, or Fab2 fragment; the binding compound is conjugated to a chemical moiety; the binding compound is in a detection 25 kit which also comprises instructions for use or disposal of reagents therein.

The invention also provides a cell which makes the antibody.

The invention embraces methods of purifying a polypeptide using a binding compound to specifically separate said polypeptides from others; of generating an antigen-binding compound complex comprising the step of contacting a sample comprising the antigen to a sample comprising a binding compound; or of modulating physiology or development of a cell expressing a receptor for a chemokine selected from TECK, MIP-3 α , or MIP-3 β ; the method comprising contacting the cell with a composition comprising an agonist or mutein of said

chemokine or an antibody antagonist of the chemokine. In certain embodiments of the method, the cell is a macrophage, lymphocyte, or eosinophil; or the physiology is a cellular calcium flux, a chemoattractant response, cellular morphology modification responses, phosphoinositide lipid turnover, or an antiviral response. In other embodiments, the receptor is DC CR, the chemokine is MIP-3α, the physiology is pulmonary physiology, or the cell is an eosinophil.

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DESCRIPTION OF THE DRAWINGS

Figures 1 and 2A-2D show chemotactic properties of mTECK recombinant protein. Fig. 1 shows migration of mouse thymocytes to recombinant mTECK and effect of pertussis toxin. Chemotaxis assays were performed as 15 described. Recombinant mouse lymphotactin was used as a positive control. Data are expressed as the mean of cell counts obtained from three separate experiments in duplicate ± SEM. In one experiment, cells were pre 20 incubated 1 h with 10 ng/ml pertussis toxin (PTX) prior to the assay. Figs. 2A-2D show migration of other leukocyte subsets to recombinant mTECK. Mouse splenic dendritic cells (Fig.2A) and mouse activated macrophages (Fig.2B) were obtained. THP-1 human monocytic cells were used without (Fig.2C) or with (Fig.2D) a 16 h activation with IFN-γ. Results are obtained as the mean of the chemotactic index from three separate experiments per cell type in duplicate ± SD. The number of cells migrating to medium alone was greater than 40 cells per 5 30 high power fields in each experiment. Recombinant MIP-1a was used as a positive control.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS I. General

The present invention provides DNA sequences encoding various mammalian proteins which exhibit structural properties characteristic of a chemotactic cytokine, or chemokine. Other embodiments are directed to chemokine receptors. See, e.g., Lodi, et al. (1994) Science 263:1762-1767; Gronenborn and Clore (1991)

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Protein Engineering 4:263-269; Miller and Kranger (1992)

Proc. Nat'l Acad. Sci. USA 89:2950-2954; Matsushima and

Oppenheim (1989) Cytokine 1:2-13; Stoeckle and Baker

(1990) New Biol. 2:313-323; Oppenheim, et al. (1991) Ann.

Rev. Immunol. 9:617-648; Schall (1991) Cytokine 3:165
183; and The Cytokine Handbook Academic Press, NY. Mouse and human embodiments are described herein.

Chemokines play an important role in immune and inflammatory responses by inducing migration and adhesion of leukocytes. These small secreted molecules are a growing superfamily of 8-14 kDa proteins characterized by a conserved four cysteine motif. See, e.g., Schall (1991) Cytokine 3:165-183; and Thomson (ed.) The Cytokine Handbook Academic Press, NY. Chemokines are secreted by activated leukocytes and act as a chemoattractant for a variety of cells which are involved in inflammation. Besides chemoattractant properties, chemokines have been shown to induce other biological responses, e.g., modulation of second messenger levels such as Ca++; inositol phosphate pool changes (see, e.g., Berridge (1993) Nature 361:315-325 or Billah and Anthes (1990) Biochem. J. 269:281-291); cellular morphology modification responses; phosphoinositide lipid turnover; possible antiviral responses; and others. Thus, the chemokines provided herein may, alone or in combination with other therapeutic reagents, have advantageous combination effects.

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Moreover, there are reasons to suggest that chemokines may have effects on other cell types, e.g., attraction or activation of monocytes, dendritic cells, T cells, eosinophils, and/or perhaps on basophils and/or neutrophils. They may also have chemoattractive effects on various neural cells including, e.g., dorsal root ganglia neurons in the peripheral nervous system and/or central nervous system neurons.

Chemokine receptors are important in the signal transduction mechanisms mediated by the chemokines. They are useful markers for distinguishing cell populations,

and have been implicated as specific receptors for retroviral infections.

The chemokine superfamily was classically divided into two groups exhibiting characteristic structural motifs, the Cys-X-Cys (C-X-C) and Cys-Cys (C-C) families. These were distinguished on the basis of a single amino acid insertion between the NH-proximal pair of cysteine residues and sequence similarity. Typically, the C-X-C chemokines, i.e., IL-8 and MGSA/Gro- α act on neutrophils 10 but not on monocytes, whereas the C-C chemokines, i.e., $\text{MIP-}1\alpha$ and RANTES, are potent chemoattractants for monocytes and lymphocytes but not neutrophils. See, e.g., Miller, et al. (1992) Crit. Rev. Immunol. 12:17-46. A recently isolated chemokine, lymphotactin, does not belong to either group and may constitute a first member 15 of a third chemokine family, the C family. Lymphotactin does not have a characteristic CC or CXC motif, and acts on lymphocytes but not neutrophils and monocytes. See, e.g., Kelner et al. (1994) <u>Science</u> 266:1395-1399. chemokine defines a new C-C chemokine family. Even more 20 recently, another chemokine exhibiting a CX3C motif has been identified, which establishes a fourth structural class.

The present invention provides additional chemokine reagents, e.g., nucleic acids, proteins and peptides, antibodies, etc., related to the newly discovered respective chemokines designated TECK; MIP-3α, and MIP-3β.

In other embodiments, the invention provides two

genes encoding novel chemokine receptors, designated DC

CR and M/DC CR. Their ligands have not yet specifically
been identified. However, the receptors exhibit

structural features typical of known chemokine receptors,
e.g., 7 transmembrane spanning structures. They may

exhibit properties of binding many different cytokines at
varying specificities (shared or promiscuous binding
specificity) or may exhibit high affinity for one
(specific) or a subset (shared) of chemokines.

The described chemokines and receptors should be important for mediating various aspects of cellular, organ, tissue, or organismal physiology or development.

5 II. Purified chemokines

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Mouse Thymus Expressed ChemoKine (TECK) nucleotide and amino acid sequences are shown in SEQ ID NO: 1 and 2, respectively. The signal sequence should run from 1 (Met) to about 23 (Ala), and removal of the signal sequence should provide one natural mature sequence beginning at 24 (Gln). Human TECK nucleotide and amino acid sequences are shown in SEQ ID NO: 3 and 4, respectively. Signal sequence cleavage is probably between about Thr and Gln.

Nucleotide and amino acid sequences of another novel chemokine, from human, designated MIP-3 α are provided in SEQ ID NO: 5 and 6, respectively. Nucleotide and derived amino acid sequences of a third novel chemokine, from human, designated MIP-3 β are shown in SEQ ID NO: 7 and 8, respectively. Signal sequence cleavage is about between Ser and Gly. Generic descriptions of physical properties of polypeptides, nucleic acids, and antibodies where directed to one embodiment clearly are generally applicable to other chemokines or receptors described herein.

The nucleotide and amino acid sequences of a novel chemokine receptor found on dendritic cells (DC), from human, and designated DC CR, are provided in SEQ ID NO: 9 and 10, respectively. The nucleotide and amino acid sequences of another novel chemokine receptor found on macrophages and dendritic cells, from human, and designated M/DC CR, are provided in SEQ ID NO: 11 and 12. Alignment of M/DC CR with the previously identified chemokine receptors CKR-1 through CKR-4 is shown in Table 1. Amino acid sequences for CKR-1 through CKR-4 are provided in SEQ ID NOS: 13-17.

The amino acid sequences for these new chemokines and receptors, provided amino to carboxy, are important in providing sequence information on the chemokine ligand

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or receptor, allowing for distinguishing the protein from other proteins. Moreover, the sequences allow preparation of peptides to generate antibodies to recognize and distinguish such segments, and allow 5 preparation of oligonucleotide probes, both of which are strategies for isolation, e.g., cloning, of genes encoding such sequences, or related sequences, e.g., natural polymorphic or other variants. Similarities of the chemokines have been observed with other cytokines.

10 See, e.g., Bosenberg, et al. (1992) Cell 71:1157-1165; Huang, et. al. (1992) Molecular Biology of the Cell 3:349-362; and Pandiella, et al. (1992) J. Biol. Chem. 267:24028-24033. Likewise for the receptors.

Table 1: Alignment of M/DC CR with CKR-1 through CKR-4. The other chemokine receptors are SEQ ID NO: 13-17. An asterisk indicates fully conserved residue among all five receptors; a period represents conservative substitutions among all five receptors.

M/DC CR MIYTRFLKGSLKMANYTLAPEDEYDVLIEGELESDEAEQCDKYDAQALS
C-C CKR-1 METPNTTEDYDTTTEFDYGDATPCQKVNERAFG
C-C CKR-2 MLSTSRSRFIRNTNESGEEVTTFFDYDYGAPCHKFDVKQIG
C-C CKR-3 MTTSLDTVETFGTTSYYDDVGLLCEKADTRALM
25 C-C CKR-4 MNPTDIADTTLDESIYSNYYLYESIPKPCTKEGIKAFG

M/DC CR
C-C CKR-1
30 C-C CKR-2
C-C CKR-3
C-C CKR-4
AQLLPPLYSLVFVIGLVGNILVVLVLVQYKRLKNMTSIYLLNLAISDLLF
AQLLPPLYSLVFIFGFVGNMLVVLILINCKKLKCLTDIYLLNLAISDLLF
AQFVPPLYSLVFTVGLLGNVVVVMILIKYRRLRIMTNIYLLNLAISDLLF
ELFLPPLYSLVFVFGLLGNSVVVLVLFKYKRLRSMTDVYLLNLAISDLLF

Table 1 (continued)

5	M/DC CR C-C CKR-1 C-C CKR-2 C-C CKR-3 C-C CKR-4	KCAFSRTPFLPADETF-WKHFLTLKMNISVLVLPLFIFTFLYVOMRKTL- TCSLHFPHESLREWKLFQALKLNLFGLVLPLLVMIICYTGIIKILI VCGPYFPRGWNNFHTIMRNILGLVLPLLIMVICYSGILKTLI LCSALYPEDTVYSWRHFHTLRMTIFCLVLPLLVMAICYTGIIKTLI YCKTKYSLNST-TWKVLSSLEINILGLVIPLGIMLFCYSMIIRTLO
10	M/DC CR C-C CKR-1 C-C CKR-2 C-C CKR-3 C-C CKR-4	RFREQRYSLFKLVFAVMVVFLLMWAPYNIAFFLSTFKEHFSLSDCKSS RRPNEKK-SKAVRLIFVIMIIFFLFWTPYNLTILISVFQDFLFTHECEQS RCRNEKKRHRAVRVIFTIMIVYFLFWTPYNIVILLNTFQEFFGLSNCEST RCPSKKK-YKAIRLIFVIMAVFFIFWTPYNVAILLSSYQSILFGNDCERS HCKNEKK-NKAVKMIFAVVVLFLGFWTPYNIVLFLETLVELEVLQDCTF
15	M/DC CR C-C CKR-1 C-C CKR-2	YNLDKSVHITKLIATTHCCINPLLYAFLDGTFSKYLCRCFH RHLDLAVQVTEVIAYTHCCVNPVIYAFVGERFRKYLRQLFH-RRVA SOLDQATQVTETLGMTHCCINPIIYAFVGEKFRSLFHIALG-CRIAPLQF
20	C-C CKR-3 C-C CKR-4	KHLDLVMLVTEVIAYSHCCMNPVIYAFVGERFRKYLRHFFH-RHLL RYLDYAIQATETLAFVHCCLNPIIYFFLGERFRKYILQLFKTCRGLFVLC
25	M/DC CR C-C CKR-1 C-C CKR-2 C-C CKR-3 C-C CKR-4	LRSNTPLQPRGQSAQGTSREEPDHSTEV*VHLVKWLPFLSVDRLERVSSTSPSTGEHELSAGF* PVCGGPGVRPGKNVKVTTQGLLDGRGKGKSIGRAPEASLQDKEGA*MHLGRYIPFLPSEKLERTSSVSPSTAEPELSIVF* QYCGLLQIYSADTPSSSYTQSTMDHDLHDAL*

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As used herein, the term "TECK" shall encompass, when used in a protein context, a protein having mature mouse or human amino acid sequences, as shown in SEQ ID NO: 2 or SEQ ID NO: 4. The invention also embraces a polypeptide comprising a significant fragment of such protein. It also refers to a polypeptide that is a species counterpart, e.g., which exhibits similar biological function, and is more homologous in natural encoding sequence than other genes from that species. Typically, such chemokine will also interact with its specific binding components, e.g., receptor. These binding components, e.g., antibodies, typically bind to the chemokine with high affinity, e.g., at least about 100 nM, usually better than about 30 nM, preferably better than about 10 nM, and more preferably at better than about 3 nM. Homologous proteins would be found in mammalian species other than mouse, e.g., rats, dogs, cats, and primates. Non-mammalian species should also possess structurally or functionally related genes and proteins.

The term "polypeptide" as used herein includes a significant fragment or segment, and encompasses a stretch of amino acid residues of at least about 8 amino acids, generally at least 10 amino acids, more generally at least 12 amino acids, often at least 14 amino acids, more often at least 16 amino acids, typically at least 18 amino acids, more typically at least 20 amino acids. usually at least 22 amino acids, more usually at least 24 amino acids, preferably at least 26 amino acids, more preferably at least 28 amino acids, and, in particularly 10 preferred embodiments, at least about 30 or more amino acids, e.g., about 35, 40, 45, 50, 60, 75, 80, 100, 120, etc. Similar proteins will likely comprise a plurality of such segments. Such fragments may have ends that begin and/or end at virtually all positions, e.g., beginning at residues 1, 2, 3, etc., and ending at, e.g., 69, 68, 67, 66, etc., in all combinations. Particularly interesting peptides have ends corresponding to structural domain boundaries.

20 The term "binding composition" refers to molecules that bind with specificity to the respective chemokine or receptor, e.g., in a ligand-receptor type fashion or an antibody-antigen interaction. These compositions may be compounds, e.g., proteins, which specifically associate with the chemokine or receptor, including natural physiologically relevant protein-protein interactions, either covalent or non-covalent. The binding composition may be a polymer, or another chemical reagent. No implication as to whether the chemokine presents a 30 concave or convex shape in its ligand-receptor interaction is represented, other than the interaction exhibit similar specificity, e.g., specific affinity. A functional analog may be a ligand with structural modifications, or may be a wholly unrelated molecule, e.g., which has a molecular shape that interacts with the appropriate ligand binding determinants. The ligands may serve as agonists or antagonists of the receptor, see, e.g., Goodman, et al. (eds.) (1990) Goodman & Gilman's:

The Pharmacological Bases of Therapeutics (8th ed.), Pergamon Press.

Substantially pure means that the protein is free from other contaminating proteins, nucleic acids, and/or 5 other biologicals typically derived from the original source organism. Purity may be assayed by standard methods, and will ordinarily be at least about 40% pure, more ordinarily at least about 50% pure, generally at least about 60% pure, more generally at least about 70% pure, often at least about 75% pure, more often at least about 80% pure, typically at least about 85% pure, more typically at least about 90% pure, preferably at least about 95% pure, more preferably at least about 98% pure, and in most preferred embodiments, at least 99% pure. Analyses will typically be by weight, but may be by molar amounts.

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Solubility of a polypeptide or fragment depends upon the environment and the polypeptide. Many parameters affect polypeptide solubility, including temperature, 20 electrolyte environment, size and molecular characteristics of the polypeptide, and nature of the solvent. Typically, the temperature at which the polypeptide is used ranges from about 4° C to about 65° C. Usually the temperature at use is greater than about 18° C and more usually greater than about 22° C. For 25 diagnostic purposes, the temperature will usually be about room temperature or warmer, but less than the denaturation temperature of components in the assay. For therapeutic purposes, the temperature will usually be body temperature, typically about 37° C for humans, 30 though under certain situations the temperature may be raised or lowered in situ or in vitro.

The electrolytes will usually approximate in situ physiological conditions, but may be modified to higher or lower ionic strength where advantageous. The actual ions may be modified, e.g., to conform to standard buffers used in physiological or analytical contexts.

The size and structure of the polypeptide should generally be in a substantially stable state, and usually not in a denatured state. The polypeptide may be associated with other polypeptides in a quaternary structure, e.g., to confer solubility, or associated with lipids or detergents in a manner that approximates natural lipid bilayer interactions.

The solvent will usually be a biologically compatible buffer, of a type used for preservation of biological activities, and will usually approximate a physiological solvent. Usually the solvent will have a neutral pH, typically between about 5 and 10, and preferably about 7.5. On some occasions, a detergent will be added, typically a mild non-denaturing one, e.g., CHS or CHAPS, or a low enough concentration as to avoid significant disruption of structural or physiological properties of the protein.

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Solubility is reflected by sedimentation measured in Svedberg units, which are a measure of the sedimentation velocity of a molecule under particular conditions. determination of the sedimentation velocity was classically performed in an analytical ultracentrifuge, but is typically now performed in a standard ultracentrifuge. See, Freifelder (1982) Physical Biochemistry (2d ed.), W.H. Freeman; and Cantor and Schimmel (1980) Biophysical Chemistry, parts 1-3, W.H. 25 Freeman & Co., San Francisco. As a crude determination, a sample containing a putatively soluble polypeptide is spun in a standard full sized ultracentrifuge at about 50K rpm for about 10 minutes, and soluble molecules will remain in the supernatant. A soluble particle or 30 polypeptide will typically be less than about 30S, more typically less than about 15S, usually less than about 10S, more usually less than about 6S, and, in particular embodiments, preferably less than about 4S, and more preferably less than about 3S.

III. Physical Variants

This invention also encompasses proteins or peptides having substantial amino acid sequence homology with the amino acid sequence of each respective chemokine or receptor. The variants include species or polymorphic variants.

Amino acid sequence homology, or sequence identity, is determined by optimizing residue matches, if necessary, by introducing gaps as required. This changes 10 when considering conservative substitutions as matches. Conservative substitutions typically include substitutions within the following groups: glycine, alanine; valine, isoleucine, leucine; aspartic acid, glutamic acid; asparagine, glutamine; serine, threonine; 15 lysine, arginine; and phenylalanine, tyrosine. Homologous amino acid sequences are typically intended to include natural allelic and interspecies variations in each respective protein sequence. Typical homologous proteins or peptides will have from 25-100% homology (if 20 gaps can be introduced), to 50-100% homology (if conservative substitutions are included) with the amino acid sequence of the appropriate chemokine or receptor. Homology measures will be at least about 35%, generally at least 40%, more generally at least 45%, often at least 50%, more often at least 55%, typically at least 60%, more typically at least 65%, usually at least 70%, more usually at least 75%, preferably at least 80%, and more preferably at least 80%, and in particularly preferred embodiments, at least 85% or more. See also Needleham, 30 et al. (1970) J. Mol. Biol. 48:443-453; Sankoff, et al. (1983) Chapter One in Time Warps. String Edits. and Macromolecules: The Theory and Practice of Sequence Comparison Addison-Wesley, Reading, MA; and software packages from IntelliGenetics, Mountain View, CA; and the University of Wisconsin Genetics Computer Group, Madison, WI.

Each of the isolated chemokine or receptor DNAs can be readily modified by nucleotide substitutions,

nucleotide deletions, nucleotide insertions, and inversions of nucleotide stretches. These modifications result in novel DNA sequences that encode these antigens, their derivatives, or proteins having similar

physiological, immunogenic, or antigenic activity. These modified sequences can be used to produce mutant antigens or to enhance expression. Enhanced expression may involve gene amplification, increased transcription, increased translation, and other mechanisms. Such mutant

chemokine or receptor derivatives include predetermined or site-specific mutations of the respective protein or its fragments. "Mutant chemokine" encompasses a polypeptide otherwise falling within the homology definition of the chemokine as set forth above, but

having an amino acid sequence that differs from that of the chemokine as found in nature, whether by way of deletion, substitution, or insertion. These include substitution levels from none, one, two, three, etc. In particular, "site specific mutant chemokine" generally

includes proteins having significant homology with a ligand having sequences of SEQ ID NO: 2, 4, 6 or 8, and as sharing various biological activities, e.g., antigenic or immunogenic, with those sequences, and in preferred embodiments contain most of the disclosed sequences.

25 Similar concepts apply to the different chemokine protein embodiments, particularly those found in various warm blooded animals, e.g., mammals and birds. As stated before, it is emphasized that descriptions are generally meant to encompass the various chemokine proteins, not limited to the mouse or human embodiments specifically discussed. Similar concepts apply to the receptor

embodiments.

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Although site specific mutation sites are often predetermined, mutants need not be site specific. Chemokine mutagenesis can be conducted by making amino acid insertions or deletions. Substitutions, deletions, insertions, or combinations may be generated to arrive at a final construct. Insertions include amino- or carboxyterminal fusions. Random mutagenesis can be conducted at

a target codon and the expressed mutants can then be screened for the desired activity. Methods for making substitution mutations at predetermined sites in DNA having a known sequence are well known in the art, e.g., by M13 primer mutagenesis or polymerase chain reaction (PCR) techniques. See also Sambrook, et al. (1989) and Ausubel, et al. (1987 and Supplements).

The mutations in the DNA normally should not place coding sequences out of reading frames and preferably will not create complementary regions that could hybridize to produce secondary mRNA structure such as loops or hairpins.

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The present invention also provides recombinant proteins, e.g., heterologous fusion proteins using segments from these proteins. A heterologous fusion protein is a fusion of proteins or segments that are naturally not normally fused in the same manner. Thus, the fusion product of an immunoglobulin with a chemokine or receptor polypeptide is a continuous protein molecule having sequences fused in a typical peptide linkage, typically made as a single translation product and exhibiting properties derived from each source peptide. A similar chimeric concept applies to heterologous nucleic acid sequences.

In addition, new constructs may be made from combining similar functional domains from other proteins. For example, ligand-binding or other segments may be "swapped" between different new fusion polypeptides or fragments. See, e.g., Cunningham, et al. (1989) Science 243:1330-1336; and O'Dowd, et al. (1988) J. Biol. Chem. 263:15985-15992. Thus, new chimeric polypeptides exhibiting new combinations of specificities will result from the functional linkage of ligand-binding specificities and other functional domains.

The phosphoramidite method described by Beaucage and Carruthers (1981) Tetra. Letts. 22:1859-1862, will produce suitable synthetic DNA fragments. A double stranded fragment will often be obtained either by synthesizing the complementary strand and annealing the

strand together under appropriate conditions or by adding the complementary strand using DNA polymerase with an appropriate primer sequence, e.g., PCR techniques.

5 IV. Functional Variants

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The blocking of physiological response to various embodiments of these chemokines may result from the inhibition of binding of the ligand to its receptor, likely through competitive inhibition. Thus, in vitro assays of the present invention will often use isolated protein, membranes from cells expressing a recombinant membrane associated chemokine, soluble fragments comprising receptor binding segments of these ligands, or fragments attached to solid phase substrates. These assays will also allow for the diagnostic determination of the effects of either binding segment mutations and modifications, or ligand mutations and modifications, e.g., ligand analogs.

This invention also contemplates the use of

competitive drug screening assays, e.g., where
neutralizing antibodies to antigen or receptor fragments
compete with a test compound for binding to the protein.
In this manner, the antibodies can be used to detect the
presence of polypeptides that share one or more antigenic
binding sites of the ligand and can also be used to
occupy binding sites on the protein that might otherwise
interact with a receptor.

Additionally, neutralizing antibodies against a specific chemokine embodiment and soluble fragments of the chemokine that contain a high affinity receptor binding site, can be used to inhibit chemokine activity in tissues, e.g., tissues experiencing abnormal physiology.

"Derivatives" of chemokine antigens include amino acid sequence mutants, glycosylation variants, and covalent or aggregate conjugates with other chemical moieties. Covalent derivatives can be prepared by linkage of functionalities to groups which are found in chemokine amino acid side chains or at the N- or C-

termini, by means which are well known in the art. These derivatives can include, without limitation, aliphatic esters or amides of the carboxyl terminus, or of residues containing carboxyl side chains, O-acyl derivatives of hydroxyl group-containing residues, and N-acyl derivatives of the amino terminal amino acid or aminogroup containing residues, e.g., lysine or arginine. Acyl groups are selected from the group of alkyl-moieties including C3 to C18 normal alkyl, thereby forming alkanoyl aroyl species. Covalent attachment to carrier proteins may be important when immunogenic moieties are haptens.

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In particular, glycosylation alterations are included, e.g., made by modifying the glycosylation patterns of a polypeptide during its synthesis and processing, or in further processing steps. Particularly preferred means for accomplishing this are by exposing the polypeptide to glycosylating enzymes derived from cells which normally provide such processing, e.g., mammalian glycosylation enzymes. Deglycosylation enzymes are also contemplated. Also embraced are versions of the same primary amino acid sequence which have other minor modifications, including phosphorylated amino acid residues, e.g., phosphotyrosine, phosphoserine, or phosphothreonine.

A major group of derivatives are covalent conjugates of the respective chemokine or receptor or fragments thereof with other proteins or polypeptides. These derivatives can be synthesized in recombinant culture such as N- or C-terminal fusions or by the use of agents known in the art for their usefulness in cross-linking proteins through reactive side groups. Preferred chemokine derivatization sites with cross-linking agents are at free amino groups, carbohydrate moieties, and 35 cysteine residues.

Fusion polypeptides between these chemokines and other homologous or heterologous proteins, e.g., other chemokines, are also provided. Many growth factors and cytokines are homodimeric entities, and a repeat

construct may have various advantages, including lessened susceptibility to proteolytic cleavage. Moreover, many receptors require dimerization to transduce a signal, and various dimeric ligands or domain repeats can be desirable. Homologous polypeptides may be fusions between different surface markers, resulting in, e.g., a hybrid protein exhibiting receptor binding specificity. Likewise, heterologous fusions may be constructed which would exhibit a combination of properties or activities of the derivative proteins. Typical examples are fusions 10 of a reporter polypeptide, e.g., luciferase, with a segment or domain of a ligand, e.g., a receptor-binding segment, so that the presence or location of the fused ligand may be easily determined. See, e.g., Dull, et al., U.S. Patent No. 4,859,609. Other gene fusion 15 partners include bacterial ß-galactosidase, trpE, Protein A, ß-lactamase, alpha amylase, alcohol dehydrogenase, a FLAG fusion, and yeast alpha mating factor. See, e.g., Godowski, et al. (1988) <u>Science</u> 241:812-816.

The phosphoramidite method described by Beaucage and Carruthers (1981) <u>Tetra. Letts.</u> 22:1859-1862, will produce suitable synthetic DNA fragments. A double stranded fragment will often be obtained either by synthesizing the complementary strand and annealing the strand together under appropriate conditions or by adding the complementary strand using DNA polymerase with an appropriate primer sequence.

Such polypeptides may also have amino acid residues that have been chemically modified by phosphorylation, sulfonation, biotinylation, or the addition or removal of other moieties, particularly those which have molecular shapes similar to phosphate groups. In some embodiments, the modifications will be useful labeling reagents, or serve as purification targets, e.g., affinity tags as FLAG.

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Fusion proteins will typically be made by either recombinant nucleic acid methods or by synthetic polypeptide methods. Techniques for nucleic acid manipulation and expression are described generally, for

example, in Sambrook, et al. (1989) Molecular Cloning: A
Laboratory Manual (2d ed.), Vols. 1-3, Cold Spring Harbor
Laboratory. Techniques for synthesis of polypeptides are
described, for example, in Merrifield (1963) J. Amer.

5 Chem. Soc. 85:2149-2156; Merrifield (1986) Science 232:
341-347; and Atherton, et al. (1989) Solid Phase Peptide
Synthesis: A Practical Approach, IRL Press, Oxford; and
chemical ligation, e.g., Dawson, et al. (1994) Science
266:776-779, a method of linking long synthetic peptides
by a peptide bond.

This invention also contemplates the use of derivatives of these chemokines or receptors other than variations in amino acid sequence or glycosylation. derivatives may involve covalent or aggregative association with chemical moieties. These derivatives generally fall into the three classes: (1) salts, (2) side chain and terminal residue covalent modifications, and (3) adsorption complexes, for example with cell membranes. Such covalent or aggregative derivatives are 20 useful as immunogens, as reagents in immunoassays, or in purification methods such as for affinity purification of ligands or other binding ligands. For example, a chemokine antigen can be immobilized by covalent bonding to a solid support such as cyanogen bromide-activated Sepharose, by methods which are well known in the art, or 25 adsorbed onto polyolefin surfaces, with or without glutaraldehyde cross-linking, for use in the assay or purification of anti-chemokine antibodies or its receptor. These chemokines can also be labeled with a detectable group, for example radioiodinated by the 30 chloramine T procedure, covalently bound to rare earth chelates, or conjugated to a fluorescent moiety for use in diagnostic assays. Purification of chemokine may be effected by immobilized antibodies or receptor.

Other modifications may be introduced with the goal of modifying the therapeutic pharmacokinetics or pharmacodynamics of a target chemokine. For example, certain means to minimize the size of the entity may

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improve its pharmacoaccessibility; other means to maximize size may affect pharmacodynamics.

A solubilized chemokine or appropriate fragment of this invention can be used as an immunogen for the production of antisera or antibodies specific for the ligand or fragments thereof. The purified chemokines can be used to screen monoclonal antibodies or chemokinebinding fragments prepared by immunization with various forms of impure preparations containing the protein. 10 particular, antibody equivalents include antigen binding fragments of natural antibodies, e.g., Fv, Fab, or F(ab)2. Purified chemokines can also be used as a reagent to detect antibodies generated in response to the presence of elevated levels of the protein or cell 15 fragments containing the protein, both of which may be diagnostic of an abnormal or specific physiological or disease condition. Additionally, chemokine protein fragments, or their concatenates, may also serve as immunogens to produce antibodies of the present invention, as described immediately below. For example, 20 this invention contemplates antibodies raised against amino acid sequences shown in SEQ ID NO: 2, 4, 6 or 8, or proteins containing them. In particular, this invention contemplates antibodies having binding affinity 25 to or being raised against specific fragments, e.g., those which are predicted to lie on the outside surfaces of protein tertiary structure. Similar concepts apply to antibodies specific for receptors of the invention.

The present invention contemplates the isolation of additional closely related species variants. Southern and Northern blot analysis should establish that similar genetic entities exist in other mammals, and establish the stringency of hybridization conditions to isolate It is likely that these chemokines and receptors 35 are widespread in species variants, e.g., rodents, lagomorphs, carnivores, artiodactyla, perissodactyla, and primates.

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The invention also provides means to isolate a group of related chemokines displaying both distinctness and

similarities in structure, expression, and function. Elucidation of many of the physiological effects of the proteins will be greatly accelerated by the isolation and characterization of distinct species variants of the ligands. In particular, the present invention provides useful probes for identifying additional homologous genetic entities in different species.

The isolated genes will allow transformation of cells lacking expression of a corresponding chemokine, e.g., either species types or cells that lack corresponding ligands and exhibit negative background activity. Expression of transformed genes will allow isolation of antigenically pure cell lines, with defined or single specie variants. This approach will allow for more sensitive detection and discrimination of the physiological effects of chemokine receptor proteins. Subcellular fragments, e.g., cytoplasts or membrane fragments, can be isolated and used.

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Dissection of critical structural elements which effect the various differentiation functions provided by ligands is possible using standard techniques of modern molecular biology, particularly in comparing members of the related class. See, e.g., the homolog-scanning mutagenesis technique described in Cunningham, et al. (1989) Science 243:1339-1336; and approaches used in O'Dowd, et al. (1988) J. Biol. Chem. 263:15985-15992; and Lechleiter, et al. (1990) EMBO J. 9:4381-4390.

In addition, receptor binding segments can be substituted between species variants to determine what structural features are important in both receptor binding affinity and specificity, as well as signal transduction. An array of different chemokine variants will be used to screen for ligands exhibiting combined properties of interaction with different receptor species variants.

Intracellular functions would probably involve segments of the receptor that are normally accessible to the cytosol. However, ligand internalization may occur under certain circumstances, and interaction between intracellular components and "extracellular" segments may occur. The specific segments of interaction of a particular chemokine with other intracellular components may be identified by mutagenesis or direct biochemical means, e.g., cross-linking or affinity methods. Structural analysis by crystallographic or other physical methods will also be applicable. Further investigation of the mechanism of signal transduction will include study of associated components that may be isolatable by affinity methods or by genetic means, e.g., complementation analysis of mutants.

Further study of the expression and control of the various chemokines will be pursued. The controlling elements associated with the proteins may exhibit differential developmental, tissue specific, or other expression patterns. Upstream or downstream genetic regions, e.g., control elements, are of interest. Differential splicing of message may lead to membrane bound forms, soluble forms, and modified versions of ligand.

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Structural studies of the proteins will lead to design of new ligands, particularly analogs exhibiting agonist or antagonist properties on the receptor. This can be combined with previously described screening methods to isolate ligands exhibiting desired spectra of activities.

Expression in other cell types will often result in glycosylation differences in a particular chemokine. Various species variants may exhibit distinct functions based upon structural differences other than amino acid sequence. Differential modifications may be responsible for differential function, and elucidation of the effects is now made possible.

Thus, the present invention provides important
reagents related to a physiological chemokine-binding protein interaction. Although the foregoing description has focused primarily upon the mouse and human embodiments of the chemokines specifically described, those of skill in the art will immediately recognize that

the invention provides other species counterparts, e.g., rat and other mammalian species or allelic or polymorphic variants.

5 V. Antibodies

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Antibodies can be raised to these chemokines, including species or polymorphic variants, and fragments thereof, both in their naturally occurring forms and in their recombinant forms. Additionally, antibodies can be raised to chemokines in either their active forms or in their inactive forms. Anti-idiotypic antibodies are also contemplated.

Antibodies, including binding fragments and single chain versions, against predetermined fragments of the ligands can be raised by immunization of animals with concatemers or conjugates of the fragments with immunogenic proteins. Monoclonal antibodies are prepared from cells secreting the desired antibody. These antibodies can be screened for binding to normal or defective chemokines, or screened for agonistic or antagonistic activity, e.g., mediated through a receptor for a chemokine. These monoclonal antibodies will usually bind with at least a KD of about 1 mM, more usually at least about 300 µM, typically at least about 10 µM, more typically at least about 30 µM, preferably at least about 10 µM, and more preferably at least about 3 µM or better.

The antibodies, including antigen binding fragments, of this invention can have significant diagnostic or therapeutic value. They can be potent antagonists that bind to ligand and inhibit binding to receptor or inhibit the ability of a ligand to elicit a biological response. They also can be useful as non-neutralizing antibodies and can be coupled to toxins or radionuclides so that when the antibody binds to ligand, a cell expressing it, e.g., on its surface via receptor, is killed. Further, these antibodies can be conjugated to drugs or other therapeutic agents, either directly or indirectly by means of a linker, and may effect drug targeting.

Antibodies to receptors may be more easily used to block ligand binding and signal transduction.

The antibodies of this invention can also be useful in diagnostic or reagent purification applications. As capture or non-neutralizing antibodies, they can be screened for ability to bind to the chemokines without inhibiting receptor binding. As neutralizing antibodies, they can be useful in competitive binding assays. They will also be useful in detecting or quantifying chemokine or, indirectly, receptors, e.g., in immunoassays. They may be used as purification reagents in immunoaffinity columns or as immunohistochemistry reagents.

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Ligand fragments may be concatenated or joined to other materials, particularly polypeptides, as fused or covalently joined polypeptides to be used as immunogens. Short peptides will preferably be made as repeat structures to increase size. A ligand and its fragments may be fused or covalently linked to a variety of immunogens, such as keyhole limpet hemocyanin, bovine serum albumin, tetanus toxoid, etc. See Microbiology, Hoeber Medical Division, Harper and Row, 1969; Landsteiner (1962) Specificity of Serological Reactions, Dover Publications, New York, and Williams, et al. (1967) Methods in Immunology and Immunochemistry, Vol. 1,

Academic Press, New York, for descriptions of methods of preparing polyclonal antisera. A typical method involves hyperimmunization of an animal with an antigen. The blood of the animal is then collected shortly after the repeated immunizations and the gamma globulin fraction is isolated.

In some instances, it is desirable to prepare monoclonal antibodies from various mammalian hosts, such as mice, rodents, primates, humans, etc. Description of techniques for preparing such monoclonal antibodies may be found in, e.g., Stites, et al. (eds.) Basic and Clinical Immunology (4th ed.), Lange Medical Publications, Los Altos, CA, and references cited therein; Harlow and Lane (1988) Antibodies: A Laboratory Manual, CSH Press; Goding (1986) Monoclonal Antibodies:

Principles and Practice (2d ed.) Academic Press, New York; and particularly in Kohler and Milstein (1975) in Nature 256:495-497, which discusses one method of generating monoclonal antibodies. Summarized briefly, 5 this method involves injecting an animal with an immunogen. The animal is then sacrificed and cells taken, e.g., from its spleen, which are then fused with myeloma cells. The result is a hybrid cell or "hybridoma" that is capable of reproducing in vitro. The population of hybridomas is then screened to isolate 10 individual clones, each of which secrete a single antibody species to the immunogen. In this manner, the individual antibody species obtained are the products of immortalized and cloned single B cells from the immune animal generated in response to a specific site 15 recognized on the immunogenic substance. Large amounts of antibody may be derived from ascites fluid from an animal.

Other suitable techniques involve in vitro exposure of lymphocytes to the antigenic polypeptides or 20 alternatively to selection of libraries of antibodies in phage or similar vectors. See, Huse, et al. (1989) "Generation of a Large Combinatorial Library of the Immunoglobulin Repertoire in Phage Lambda, " Science 246:1275-1281; and Ward, et al. (1989) Nature 341:544-25 546. The polypeptides and antibodies of the present invention may be used with or without modification, including chimeric or humanized antibodies. Frequently, the polypeptides and antibodies will be labeled by joining, either covalently or non-covalently, a substance 30 which provides for a detectable signal. A wide variety of labels and conjugation techniques are known and are reported extensively in both the scientific and patent literature. Suitable labels include radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent moieties, chemiluminescent moieties, magnetic particles, and the like. Patents, teaching the use of such labels include U.S. Patent Nos. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241. Also,

recombinant immunoglobulins may be produced, see Cabilly, U.S. Patent No. 4,816,567; and Queen et al. (1989) <u>Proc.</u> Nat'l. Acad. Sci. 86:10029-10033.

The antibodies of this invention can also be used for affinity chromatography in isolating the protein. Columns can be prepared where the antibodies are linked to a solid support, e.g., particles, such as agarose, Sephadex, or the like, where a cell lysate may be passed through the column, the column washed, followed by increasing concentrations of a mild denaturant, whereby the purified chemokine protein will be released.

The antibodies may also be used to screen expression libraries for particular expression products. Usually the antibodies used in such a procedure will be labeled with a moiety allowing easy detection of presence of antigen by antibody binding.

Antibodies raised against these chemokine will also be useful to raise anti-idiotypic antibodies. These will be useful in detecting or diagnosing various immunological conditions related to expression of the respective antigens.

VI. Nucleic Acids

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reagents are useful in isolating a DNA clone encoding these chemokines, e.g., from a natural source.

Typically, it will be useful in isolating a gene from another individual, and similar procedures will be applied to isolate genes from other species, e.g., warm blooded animals, such as birds and mammals. Cross hybridization will allow isolation of ligand from other species. A number of different approaches should be available to successfully isolate a suitable nucleic acid clone. Similar concepts apply to the receptor embodiments.

The purified protein or defined peptides are useful for generating antibodies by standard methods, as described above. Synthetic peptides or purified protein can be presented to an immune system to generate

monoclonal or polyclonal antibodies. See, e.g., Coligan (1991) <u>Current Protocols in Immunology</u> Wiley/Greene; and Harlow and Lane (1989) <u>Antibodies: A Laboratory Manual</u> Cold Spring Harbor Press. Alternatively, a chemokine receptor can be used as a specific binding reagent, and advantage can be taken of its specificity of binding, much like an antibody would be used. However, chemokine receptors are typically 7 transmembrane proteins, which could be sensitive to appropriate interaction with lipid or membrane. The signal transduction typically is mediated through a G-protein.

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For example, the specific binding composition could be used for screening of an expression library made from a cell line which expresses a particular chemokine. The screening can be standard staining of surface expressed ligand, or by panning. Screening of intracellular expression can also be performed by various staining or immunofluorescence procedures. The binding compositions could be used to affinity purify or sort out cells expressing the ligand.

The peptide segments can also be used to predict appropriate oligonucleotides to screen a library, e.g., to isolate species variants. The genetic code can be used to select appropriate oligonucleotides useful as probes for screening. In combination with polymerase chain reaction (PCR) techniques, synthetic oligonucleotides will be useful in selecting correct clones from a library. Complementary sequences will also be used as probes or primers. Based upon identification of the likely amino terminus, the third peptide should be particularly useful, e.g., coupled with anchored vector or poly-A complementary PCR techniques or with complementary DNA of other peptides.

This invention contemplates use of isolated DNA or fragments to encode a biologically active corresponding chemokine polypeptide. In addition, this invention covers isolated or recombinant DNA which encodes a biologically active protein or polypeptide which is capable of hybridizing under appropriate conditions with

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the DNA sequences described herein. Said biologically active protein or polypeptide can be an intact ligand, or fragment, and have an amino acid sequence as disclosed in SEQ ID NOS: 1, 3, 5 and 7. Further, this invention covers the use of isolated or recombinant DNA, or fragments thereof, which encode proteins which are homologous to a chemokine or which was isolated using cDNA encoding a chemokine as a probe. The isolated DNA can have the respective regulatory sequences in the 5' and 3' flanks, e.g., promoters, enhancers, poly-A addition signals, and others.

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An "isolated" nucleic acid is a nucleic acid, e.g., an RNA, DNA, or a mixed polymer, which is substantially separated from other components which naturally accompany a native sequence, e.g., ribosomes, polymerases, and flanking genomic sequences from the originating species. The term embraces a nucleic acid sequence which has been removed from its naturally occurring environment, and includes recombinant or cloned DNA isolates and chemically synthesized analogs or analogs biologically synthesized by heterologous systems. A substantially pure molecule includes isolated forms of the molecule.

An isolated nucleic acid will generally be a homogeneous composition of molecules, but will, in some embodiments, contain minor heterogeneity. This heterogeneity is typically found at the polymer ends or portions not critical to a desired biological function or activity.

A "recombinant" nucleic acid is defined either by its method of production or its structure. In reference to its method of production, e.g., a product made by a process, the process is use of recombinant nucleic acid techniques, e.g., involving human intervention in the nucleotide sequence, typically selection or production. 35 Alternatively, it can be a nucleic acid made by generating a sequence comprising fusion of two fragments which are not naturally contiguous to each other, but is meant to exclude products of nature, e.g., naturally occurring mutants. Thus, for example, products made by

transforming cells with any unnaturally occurring vector is encompassed, as are nucleic acids comprising sequence derived using any synthetic oligonucleotide process. Such is often done to replace a codon with a redundant codon encoding the same or a conservative amino acid, while typically introducing or removing a sequence recognition site. Alternatively, it is performed to join together nucleic acid segments of desired functions to generate a single genetic entity comprising a desired combination of functions not found in the commonly available natural forms. Restriction enzyme recognition sites are often the target of such artificial manipulations, but other site specific targets, e.g., promoters, DNA replication sites, regulation sequences, 15 control sequences, or other useful features may be incorporated by design. A similar concept is intended for a recombinant, e.g., fusion, polypeptide. Specifically included are synthetic nucleic acids which, by genetic code redundancy, encode polypeptides similar to fragments of these antigens, and fusions of sequences from various different species variants.

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A significant "fragment" in a nucleic acid context is a contiguous segment of at least about 17 nucleotides, generally at least about 20 nucleotides, more generally at least about 23 nucleotides, ordinarily at least about 26 nucleotides, more ordinarily at least about 29 nucleotides, often at least about 32 nucleotides, more often at least about 35 nucleotides, typically at least about 38 nucleotides, more typically at least about 41 nucleotides, usually at least about 44 nucleotides, more usually at least about 47 nucleotides, preferably at least about 50 nucleotides, more preferably at least about 53 nucleotides, and in particularly preferred embodiments will be at least about 56 or more 35 nucleotides, e.g., 60, 65, 75, 85, 100, 120, 150, 200, 250, 300, 400, etc. Such fragments may have ends which begin and/or end at virtually all positions, e.g., beginning at nucleotides 1, 2, 3, etc., and ending at, e.g., 300, 299, 298, 287, etc., in all combinations.

Particularly interesting polynucleotides have ends corresponding to structural domain boundaries.

A DNA which codes for a particular chemokine protein or peptide will be very useful to identify genes, mRNA, and cDNA species which code for related or homologous ligands, as well as DNAs which code for homologous proteins from different species. There are likely homologs in other species, including primates. Various chemokine proteins should be homologous and are encompassed herein. However, even proteins that have a more distant evolutionary relationship to the ligand can readily be isolated under appropriate conditions using these sequences if they are sufficiently homologous. Primate chemokines are of particular interest.

This invention further covers recombinant DNA molecules and fragments having a DNA sequence identical to or highly homologous to the isolated DNAs set forth herein. In particular, the sequences will often be operably linked to DNA segments which control transcription, translation, and DNA replication. Alternatively, recombinant clones derived from the genomic sequences, e.g., containing introns, will be useful for transgenic studies, including, e.g., transgenic cells and organisms, and for gene therapy.

See, e.g., Goodnow (1992) "Transgenic Animals" in Roitt

See, e.g., Goodnow (1992) "Transgenic Animals" in Roitt (ed.) Encyclopedia of Immunology Academic Press, San Diego, pp. 1502-1504; Travis (1992) Science 256:1392-1394; Kuhn, et al. (1991) Science 254:707-710; Capecchi (1989) Science 244:1288; Robertson (1987) (ed.)

Teratocarcinomas and Embryonic Stem Cells: A Practical Approach IRL Press, Oxford; and Rosenberg (1992) J. Clinical Oncology 10:180-199.

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Homologous nucleic acid sequences, when compared, exhibit significant similarity, or identity. The standards for homology in nucleic acids are either measures for homology generally used in the art by sequence comparison or based upon hybridization conditions. The hybridization conditions are described in greater detail below.

Substantial homology in the nucleic acid sequence comparison context means either that the segments, or their complementary strands, when compared, are identical when optimally aligned, with appropriate nucleotide insertions or deletions, in at least about 50% of the nucleotides, generally at least about 56%, more generally at least about 59%, ordinarily at least about 62%, more ordinarily at least about 65%, often at least about 68%, more often at least about 71%, typically at least about 74%, more typically at least about 77%, usually at least about 80%, more usually at least about 85%, preferably at least about 90%, more preferably at least about 95 to 98% or more, and in particular embodiments, as high at about 99% or more of the nucleotides. Alternatively, substantial homology exists when the segments will hybridize under selective hybridization conditions, to a strand, or its complement, typically using a sequence derived from those set forth in SEQ ID NOS: 1, 3, 4, 7, 9 or 11. Typically, selective hybridization will occur when there is at least about 55% homology over a stretch 20 of at least about 30 nucleotides, preferably at least about 65% over a stretch of at least about 25 nucleotides, more preferably at least about 75%, and most

See, Kanehisa (1984) Nuc. Acids Res. 12:203-213. The length of homology comparison, as described, may be over longer stretches, and in certain embodiments will be over a stretch of at least about 17 nucleotides, usually at least about 20 nucleotides, more usually at least about 24 nucleotides, typically at least about 28 nucleotides, more typically at least about 40 nucleotides, preferably at least about 50 nucleotides, and more preferably at

least about 75 to 100 or more nucleotides.

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preferably at least about 90% over about 20 nucleotides.

Stringent conditions, in referring to homology in the hybridization context, will be stringent combined conditions of salt, temperature, organic solvents, and other parameters, typically those controlled in hybridization reactions. Stringent temperature conditions will usually include temperatures in excess of about 30° C, more usually in excess of about 37° C, typically in excess of about 45° C, more typically in excess of about 55° C, preferably in excess of about 65° C, and more preferably in excess of about 70° C.

5 Stringent salt conditions will ordinarily be less than about 1000 mM, usually less than about 500 mM, more usually less than about 400 mM, typically less than about 300 mM, preferably less than about 200 mM, and more preferably less than about 150 mM. However, the combination of parameters is much more important than the measure of any single parameter. See, e.g., Wetmur and

Davidson (1968) J. Mol. Biol. 31:349-370.

Corresponding chemokines from other mammalian species can be cloned and isolated by cross-species hybridization of closely related species. Alternatively, sequences from a data base may be recognized as having similarity. Homology may be relatively low between distantly related species, and thus hybridization of relatively closely related species is advisable.

20 Alternatively, preparation of an antibody preparation which exhibits less species specificity may be useful in

expression cloning approaches. PCR approaches using segments of conserved sequences will also be used.

25 VII. Making chemokine; Mimetics

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DNA which encodes each respective chemokine or fragments thereof can be obtained by chemical synthesis, screening cDNA libraries, or by screening genomic libraries prepared from a wide variety of cell lines or tissue samples.

This DNA can be expressed in a wide variety of host cells for the synthesis of a full-length ligand or fragments which can in turn, for example, be used to generate polyclonal or monoclonal antibodies; for binding studies; for construction and expression of modified molecules; and for structure/function studies. Each antigen or its fragments can be expressed in host cells that are transformed or transfected with appropriate expression vectors. These molecules can be substantially

purified to be free of protein or cellular contaminants, other than those derived from the recombinant host, and therefore are particularly useful in pharmaceutical compositions when combined with a pharmaceutically acceptable carrier and/or diluent. The antigen, or portions thereof, may be expressed as fusions with other proteins.

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Expression vectors are typically self-replicating DNA or RNA constructs containing the desired antigen gene or its fragments, usually operably linked to suitable genetic control elements that are recognized in a suitable host cell. These control elements are capable of effecting expression within a suitable host. specific type of control elements necessary to effect expression will depend upon the eventual host cell used. Generally, the genetic control elements can include a prokaryotic promoter system or a eukaryotic promoter expression control system, and typically include a transcriptional promoter, an optional operator to control the onset of transcription, transcription enhancers to elevate the level of mRNA expression, a sequence that encodes a suitable ribosome binding site, and sequences that terminate transcription and translation. Expression vectors also usually contain an origin of replication that allows the vector to replicate independently of the host cell.

The vectors of this invention contain DNA which encodes embodiments of a chemokine, receptor, or a fragment thereof, typically encoding a biologically active polypeptide. The DNA can be under the control of a viral promoter and can encode a selection marker. This invention further contemplates use of such expression vectors which are capable of expressing eukaryotic cDNA coding for each chemokine in a prokaryotic or eukaryotic host, where the vector is compatible with the host and where the eukaryotic cDNA coding for the ligand is inserted into the vector such that growth of the host containing the vector expresses the cDNA in question. Usually, expression vectors are designed for stable

replication in their host cells or for amplification to greatly increase the total number of copies of the desirable gene per cell. It is not always necessary to require that an expression vector replicate in a host cell, e.g., it is possible to effect transient expression of the ligand or its fragments in various hosts using vectors that do not contain a replication origin that is recognized by the host cell. It is also possible to use vectors that cause integration of a chemokine gene or its fragments into the host DNA by recombination, or to integrate a promoter which controls expression of an endogenous gene.

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Vectors, as used herein, comprise plasmids, viruses, bacteriophage, integratable DNA fragments, and other

vehicles which enable the integration of DNA fragments into the genome of the host. Expression vectors are specialized vectors which contain genetic control elements that effect expression of operably linked genes. Plasmids are the most commonly used form of vector but all other forms of vectors which serve an equivalent function and which are, or become, known in the art are suitable for use herein. See, e.g., Pouwels, et al. (1985 and Supplements) Cloning Vectors: A Laboratory Manual, Elsevier, N.Y., and Rodriquez, et al. (1988) (eds.) Vectors: A Survey of Molecular Cloning

<u>Vectors and Their Uses</u>, Buttersworth, Boston, MA.

Transformed cells include cells, preferably mammalian, that have been transformed or transfected with a chemokine gene containing vector constructed using recombinant DNA techniques. Transformed host cells usually express the ligand or its fragments, but for purposes of cloning, amplifying, and manipulating its DNA, do not need to express the protein. This invention further contemplates culturing transformed cells in a nutrient medium, thus permitting the protein to accumulate in the culture. The protein can be recovered, either from the culture or from the culture medium.

For purposes of this invention, DNA sequences are operably linked when they are functionally related to

each other. For example, DNA for a presequence or secretory leader is operably linked to a polypeptide if it is expressed as a preprotein or participates in directing the polypeptide to the cell membrane or in secretion of the polypeptide. A promoter is operably linked to a coding sequence if it controls the transcription of the polypeptide; a ribosome binding site is operably linked to a coding sequence if it is positioned to permit translation. Usually, operably linked means contiguous and in reading frame, however, certain genetic elements such as repressor genes are not contiguously linked but still bind to operator sequences that in turn control expression.

Suitable host cells include prokaryotes, lower

15 eukaryotes, and higher eukaryotes. Prokaryotes include
both gram negative and gram positive organisms, e.g., E.
coli and B. subtilis. Lower eukaryotes include yeasts,
e.g., S. cerevisiae and Pichia, and species of the genus
Dictyostelium. Higher eukaryotes include established

20 tissue culture cell lines from animal cells, both of
non-mammalian origin, e.g., insect cells, and birds, and
of mammalian origin, e.g., human, primates, and rodents.

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Prokaryotic host-vector systems include a wide variety of vectors for many different species. As used herein, E. coli and its vectors will be used generically to include equivalent vectors used in other prokaryotes. A representative vector for amplifying DNA is pBR322 or many of its derivatives. Vectors that can be used to express these chemokines or their fragments include, but are not limited to, such vectors as those containing the lac promoter (pUC-series); trp promoter (pBR322-trp); Ipp promoter (the pIN-series); lambda-pP or pR promoters (pOTS); or hybrid promoters such as ptac (pDR540). See Brosius, et al. (1988) "Expression Vectors Employing Lambda-, trp-, lac-, and Ipp-derived Promoters," in Rodriguez and Denhardt (eds.) Vectors: A Survey of Molecular Cloning Vectors and Their Uses, Buttersworth, Boston, Chapter 10, pp. 205-236.

Lower eukaryotes, e.g., yeasts and Dictyostelium, may be transformed with chemokine sequence containing vectors. For purposes of this invention, the most common lower eukaryotic host is the baker's yeast, Saccharomyces 5 cerevisiae. It will be used to generically represent lower eukaryotes although a number of other strains and species are also available. Yeast vectors typically consist of a replication origin (unless of the integrating type), a selection gene, a promoter, DNA 10 encoding the desired protein or its fragments, and sequences for translation termination, polyadenylation, and transcription termination. Suitable expression vectors for yeast include such constitutive promoters as 3-phosphoglycerate kinase and various other glycolytic 15 enzyme gene promoters or such inducible promoters as the alcohol dehydrogenase 2 promoter or metallothionine promoter. Suitable vectors include derivatives of the following types: self-replicating low copy number (such as the YRp-series), self-replicating high copy number (such as the YEp-series); integrating types (such as the YIp-series), or mini-chromosomes (such as the YCpseries).

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Higher eukaryotic tissue culture cells are the preferred host cells for expression of the functionally active chemokine protein. In principle, most any higher eukaryotic tissue culture cell line is workable, e.g., insect baculovirus expression systems, whether from an invertebrate or vertebrate source. However, mammalian cells are preferred, in that the processing, both 30 cotranslationally and posttranslationally. Transformation or transfection and propagation of such cells has become a routine procedure. Examples of useful cell lines include HeLa cells, Chinese hamster ovary (CHO) cell lines, baby rat kidney (BRK) cell lines, . 35 insect cell lines, bird cell lines, and monkey (COS) cell lines. Expression vectors for such cell lines usually include an origin of replication, a promoter, a translation initiation site, RNA splice sites (if genomic DNA is used), a polyadenylation site, and a transcription termination site. These vectors also usually contain a selection gene or amplification gene. Suitable expression vectors may be plasmids, viruses, or retroviruses carrying promoters derived, e.g., from such sources as from adenovirus, SV40, parvoviruses, vaccinia virus, or cytomegalovirus. Representative examples of suitable expression vectors include pCDNA1; pCD, see Okayama, et al. (1985) Mol. Cell Biol. 5:1136-1142; pMClneo Poly-A, see Thomas, et al. (1987) Cell 51:503-512; and a baculovirus vector such as pAC 373 or pAC 610.

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It will often be desired to express a chemokine polypeptide in a system which provides a specific or defined glycosylation pattern. In this case, the usual pattern will be that provided naturally by the expression system. However, the pattern will be modifiable by exposing the polypeptide, e.g., an unglycosylated form, to appropriate glycosylating proteins introduced into a heterologous expression system. For example, a chemokine gene may be co-transformed with one or more genes encoding mammalian or other glycosylating enzymes. Using this approach, certain mammalian glycosylation patterns will be achievable or approximated in prokaryote or other cells.

A chemokine, or a fragment thereof, may be engineered to be phosphatidyl inositol (PI) linked to a cell membrane, but can be removed from membranes by treatment with a phosphatidyl inositol cleaving enzyme, e.g., phosphatidyl inositol phospholipase-C. This releases the antigen in a biologically active form, and allows purification by standard procedures of protein chemistry. See, e.g., Low (1989) <u>Biochim. Biophys. Acta 988:427-454</u>; Tse, et al. (1985) <u>Science 230:1003-1008</u>; and Brunner, et al. (1991) <u>J. Cell Biol.</u> 114:1275-1283.

Now that these chemokines have been characterized, fragments or derivatives thereof can be prepared by conventional processes for synthesizing peptides. These include processes such as are described in Stewart and Young (1984) Solid Phase Peptide Synthesis, Pierce Chemical Co., Rockford, IL; Bodanszky and Bodanszky

(1984) The Practice of Peptide Synthesis, Springer-Verlag; New York; and Bodanszky (1984) The Principles of Peptide Synthesis, Springer-Verlag, New York. For example, an azide process, an acid chloride process, an acid anhydride process, an acid anhydride process, an active ester process (for example, p-nitrophenyl ester, N-hydroxysuccinimide ester, or cyanomethyl ester), a carbodiimidazole process, an oxidative-reductive process, or a dicyclohexylcarbodiimide (DCCD)/additive process can be used. Solid phase and solution phase syntheses are both applicable to the foregoing processes.

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These chemokines, fragments, or derivatives are suitably prepared in accordance with the above processes as typically employed in peptide synthesis, generally either by a so-called stepwise process which comprises condensing an amino acid to the terminal amino acid, one by one in sequence, or by coupling peptide fragments to the terminal amino acid. Amino groups that are not being used in the coupling reaction are typically protected to prevent coupling at an incorrect location.

If a solid phase synthesis is adopted, the C-terminal amino acid is typically bound to an insoluble carrier or support through its carboxyl group. The insoluble carrier is not particularly limited as long as it has a binding capability to a reactive carboxyl group. Examples of such insoluble carriers include halomethyl resins, such as chloromethyl resin or bromomethyl resin, hydroxymethyl resins, phenol resins, tertalkyloxycarbonyl-hydrazidated resins, and the like.

An amino group-protected amino acid is bound in sequence through condensation of its activated carboxyl group and the reactive amino group of the previously formed peptide or chain, to synthesize the peptide step by step. After synthesizing the complete sequence, the peptide is split off from the insoluble carrier to produce the peptide. This solid-phase approach is generally described by Merrifield, et al. (1963) in J. Am. Chem. Soc. 85:2149-2156.

The prepared ligand and fragments thereof can be isolated and purified from the reaction mixture by means of peptide separation, e.g., by extraction, precipitation, electrophoresis and various forms of chromatography, and the like. The various chemokines of this invention can be obtained in varying degrees of purity depending upon its desired use. Purification can be accomplished by use of the protein purification techniques disclosed herein or by the use of the antibodies herein described, e.g., in immunoabsorbant 10 affinity chromatography. This immunoabsorbant affinity chromatography is carried out by first linking the antibodies to a solid support and then contacting the linked antibodies with solubilized lysates of appropriate source cells, lysates of other cells expressing the ligand, or lysates or supernatants of cells producing the desired chemokine as a result of DNA techniques, see below.

20 VIII. Uses

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The present invention provides reagents which will find use in diagnostic applications as described elsewhere herein, e.g., in the general description for developmental abnormalities, or below in the description of kits for diagnosis.

This invention also provides reagents with significant therapeutic value. These chemokines (naturally occurring or recombinant), fragments thereof and antibodies thereto, along with compounds identified as having binding affinity to them, should be useful in the treatment of conditions associated with abnormal physiology or development, including inflammatory conditions, including asthma. In particular, modulation of trafficking of leukocytes is one likely biological activity, but a wider tissue distribution might suggest broader biological activity, including, e.g., antiviral effects. Abnormal proliferation, regeneration, degeneration, and atrophy may be modulated by appropriate therapeutic treatment using the compositions provided

herein. For example, a disease or disorder associated with abnormal expression or abnormal signaling by a chemokine should be a likely target for an agonist or antagonist of the ligand.

Various abnormal physiological or developmental conditions are known in cell types shown to possess the chemokine mRNAs by Northern blot analysis. See Berkow (ed.) The Merck Manual of Diagnosis and Therapy, Merck & Co., Rahway, N.J.; and Thorn, et al. Harrison's

Principles of Internal Medicine, McGraw-Hill, N.Y.

Developmental or functional abnormalities, e.g., of the immune system, cause significant medical abnormalities and conditions which may be susceptible to prevention or treatment using compositions provided herein.

15 Chemokine antibodies, including recombinant forms, can be purified and then administered to a patient. These reagents can be combined for therapeutic use with additional active or inert ingredients, e.g., in conventional pharmaceutically acceptable carriers or 20 diluents, e.g., immunogenic adjuvants, along with physiologically innocuous stabilizers and excipients. These combinations can be sterile filtered and placed into dosage forms as by lyophilization in dosage vials or storage in stabilized aqueous preparations. This 25 invention also contemplates use of antibodies or binding fragments thereof, including forms which are not complement binding. Moreover, modifications to the antibody molecules or antigen binding fragments thereof, may be adopted which affect the pharmacokinetics or pharmacodynamics of the therapeutic entity. 30

Drug screening using antibodies or receptor or fragments thereof can be performed to identify compounds having binding affinity to each chemokine or receptor, including isolation of associated components. Subsequent biological assays can then be utilized to determine if the compound has intrinsic stimulating activity and is therefore a blocker or antagonist in that it blocks the activity of the ligand. Likewise, a compound having intrinsic stimulating activity can activate the receptor

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and is thus an agonist in that it simulates the activity of a chemokine. This invention further contemplates the therapeutic use of antibodies to these chemokines as antagonists. This approach should be particularly useful with other chemokine species variants.

The quantities of reagents necessary for effective therapy will depend upon many different factors, including means of administration, target site, physiological state of the patient, and other medicants administered. Thus, treatment dosages should be titrated 10 to optimize safety and efficacy in various populations, including racial subgroups, age, gender, etc. Typically, dosages used in vitro may provide useful guidance in the amounts useful for in situ administration of these 15 reagents. Animal testing of effective doses for treatment of particular disorders will provide further predictive indication of human dosage. Various considerations are described, e.g., in Gilman, et al. (eds.) (1990) Goodman and Gilman's: The Pharmacological Bases of Therapeutics, 8th Ed., Pergamon Press; and 20 Remington's Pharmaceutical Sciences, 17th ed. (1990), Mack Publishing Co., Easton, Penn.. Methods for administration are discussed therein and below, e.g., for oral, intravenous, intraperitoneal, or intramuscular administration, transdermal diffusion, and others. 25 Pharmaceutically acceptable carriers typically include water, saline, buffers, and other compounds described, e.g., in the Merck Index, Merck & Co., Rahway, New Jersey. Dosage ranges would ordinarily be expected to be in amounts lower than 1 mM concentrations, typically less 30 than about 10 µM concentrations, usually less than about 100 nM, preferably less than about 10 pM (picomolar), and most preferably less than about 1 fM (femtomolar), with an appropriate carrier. Slow release formulations, or a slow release apparatus will often be utilized for 35 continuous administration.

A chemokine, fragments thereof, or antibodies to it or its fragments, antagonists, and agonists, may be administered directly to the host to be treated or,

depending on the size of the compounds, it may be desirable to conjugate them to carrier proteins such as ovalbumin or serum albumin prior to their administration. Therapeutic formulations may be administered in any conventional dosage formulation. While it is possible for the active ingredient to be administered alone, it is often preferable to present it as a pharmaceutical formulation. Formulations typically comprise at least one active ingredient, as defined above, together with 10 one or more acceptable carriers thereof. Each carrier should be both pharmaceutically and physiologically acceptable in the sense of being compatible with the other ingredients and not injurious to the patient. Carriers may improve storage life, stability, etc. Formulations include those suitable for oral, rectal, 15 nasal, or parenteral (including subcutaneous, intramuscular, intravenous and intradermal) administration. The formulations may conveniently be presented in unit dosage form and may be prepared by any methods well known in the art of pharmacy. See, e.g., 20 Gilman, et al. (eds.) (1990) Goodman and Gilman's: The Pharmacological Bases of Therapeutics, 8th Ed., Pergamon Press; and Remington's Pharmaceutical Sciences, 17th ed. (1990), Mack Publishing Co., Easton, Penn.; Avis, et al. (eds.) (1993) Pharmaceutical Dosage Forms: Parenteral Medications Dekker, New York; Lieberman, et al. (eds.) (1990) Pharmaceutical Dosage Forms: Tablets Dekker, New York; and Lieberman, et al. (eds.) (1990) Pharmaceutical Dosage Forms: Disperse Systems Dekker, New York. The therapy of this invention may be combined with or used in 30 association with other therapeutic agents.

Both the naturally occurring and the recombinant forms of the chemokines of this invention are particularly useful in kits and assay methods which are capable of screening compounds for binding activity to the proteins. Several methods of automating assays have been developed in recent years so as to permit screening of tens of thousands of compounds in a short period. See, e.g., Fodor, et al. (1991) Science 251:767-773,

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which describes means for testing of binding affinity by a plurality of defined polymers synthesized on a solid substrate. The development of suitable assays can be greatly facilitated by the availability of large amounts of purified, soluble chemokine as provided by this invention.

For example, antagonists can normally be found once the ligand has been structurally defined. Testing of potential ligand analogs is now possible upon the development of highly automated assay methods using physiologically responsive cells. In particular, new agonists and antagonists will be discovered by using screening techniques described herein.

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Viable cells could also be used to screen for the effects of drugs on respective chemokine mediated 15 functions, e.g., second messenger levels, i.e., Ca++; inositol phosphate pool changes (see, e.g., Berridge (1993) Nature 361:315-325 or Billah and Anthes (1990) Biochem. J. 269:281-291); cellular morphology modification responses; phosphoinositide lipid turnover; 20 an antiviral response, and others. Some detection methods allow for elimination of a separation step, e.g., a proximity sensitive detection system. Calcium sensitive dyes will be useful for detecting Ca++ levels, with a fluorimeter or a fluorescence cell sorting 25 apparatus.

Rational drug design may also be based upon structural studies of the molecular shapes of the chemokines and other effectors or analogs. Effectors may be other proteins which mediate other functions in response to ligand binding, or other proteins which normally interact with the receptor. One means for determining which sites interact with specific other proteins is a physical structure determination, e.g., x-35 ray crystallography or 2 dimensional NMR techniques. These will provide guidance as to which amino acid residues form molecular contact regions. For a detailed description of protein structural determination, see,

e.g., Blundell and Johnson (1976) <u>Protein</u> <u>Crystallography</u>, Academic Press, New York.

Purified chemokine can be coated directly onto plates for use in the aforementioned drug screening techniques. However, non-neutralizing antibodies to these ligands can be used as capture antibodies to immobilize the respective ligand on the solid phase.

Similar concepts also apply to the chemokine receptor embodiments of the invention.

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IX. Kits

This invention also contemplates use of chemokine proteins, fragments thereof, peptides, and their fusion products in a variety of diagnostic kits and methods for detecting the presence of ligand, antibodies, or chemokine receptors. Typically the kit will have a compartment containing either a defined chemokine peptide or gene segment or a reagent which recognizes one or the other, e.g., antibodies.

20 A kit for determining the binding affinity of a test compound to a chemokine would typically comprise a test compound; a labeled compound, for example an antibody having known binding affinity for the ligand; a source of chemokine (naturally occurring or recombinant); and a means for separating bound from free labeled compound, 25 such as a solid phase for immobilizing the ligand. Once compounds are screened, those having suitable binding affinity to the ligand can be evaluated in suitable biological assays, as are well known in the art, to 30 determine whether they act as agonists or antagonists to the receptor. The availability of recombinant chemokine polypeptides also provides well-defined standards for calibrating such assays or as positive control samples.

A preferred kit for determining the concentration
of, for example, a chemokine in a sample would typically
comprise a labeled compound, e.g., antibody, having known
binding affinity for the ligand, a source of ligand
(naturally occurring or recombinant) and a means for
separating the bound from free labeled compound, for

example, a solid phase for immobilizing the chemokine. Compartments containing reagents, and instructions for use or disposal, will normally be provided.

Antibodies, including antigen binding fragments, specific for the chemokine or ligand fragments are useful in diagnostic applications to detect the presence of elevated levels of chemokine and/or its fragments. Such diagnostic assays can employ lysates, live cells, fixed cells, immunofluorescence, cell cultures, body fluids, and further can involve the detection of antigens related to the ligand in serum, or the like. Diagnostic assays may be homogeneous (without a separation step between free reagent and antigen-ligand complex) or heterogeneous (with a separation step). Various commercial assays exist, such as radioimmunoassay (RIA), enzyme-linked 15 immunosorbent assay (ELISA), enzyme immunoassay (EIA), enzyme-multiplied immunoassay technique (EMIT), substrate-labeled fluorescent immunoassay (SLFIA), and the like. For example, unlabeled antibodies can be 20 employed by using a second antibody which is labeled and which recognizes the antibody to a chemokine or to a particular fragment thereof. Similar assays have also been extensively discussed in the literature. See, e.g., Harlow and Lane (1988) Antibodies: A Laboratory Manual. CSH.

Anti-idiotypic antibodies may have similar uses to diagnose presence of antibodies against a chemokine, as such may be diagnostic of various abnormal states. For example, overproduction of a chemokine may result in production of various immunological reactions which may be diagnostic of abnormal physiological states, particularly in various inflammatory or asthma conditions.

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Frequently, the reagents for diagnostic assays are supplied in kits, so as to optimize the sensitivity of the assay. For the subject invention, depending upon the nature of the assay, the protocol, and the label, either labeled or unlabeled antibody or labeled chemokine is provided. This is usually in conjunction with other

additives, such as buffers, stabilizers, materials necessary for signal production such as substrates for enzymes, and the like. Preferably, the kit will also contain instructions for proper use and disposal of the contents after use. Typically the kit has compartments for each useful reagent. Desirably, the reagents are provided as a dry lyophilized powder, where the reagents may be reconstituted in an aqueous medium providing appropriate concentrations of reagents for performing the assay.

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The aforementioned constituents of the drug screening and the diagnostic assays may be used without modification or may be modified in a variety of ways. For example, labeling may be achieved by covalently or non-covalently joining a moiety which directly or 15 indirectly provides a detectable signal. In any of these assays, the ligand, test compound, chemokine, or antibodies thereto can be labeled either directly or indirectly. Possibilities for direct labeling include label groups: radiolabels such as 125I, enzymes (U.S. 20 Pat. No. 3,645,090) such as peroxidase and alkaline phosphatase, and fluorescent labels (U.S. Pat. No. 3,940,475) capable of monitoring the change in fluorescence intensity, wavelength shift, or fluorescence polarization. Possibilities for indirect labeling 25 include biotinylation of one constituent followed by binding to avidin coupled to one of the above label groups.

There are also numerous methods of separating bound from the free ligand, or alternatively bound from free test compound. The chemokine can be immobilized on various matrixes followed by washing. Suitable matrixes include plastic such as an ELISA plate, filters, and beads. Methods of immobilizing the chemokine to a matrix include, without limitation, direct adhesion to plastic, use of a capture antibody, chemical coupling, and biotinavidin. The last step in this approach involves the precipitation of ligand/antibody complex by any of several methods including those utilizing, e.g., an

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organic solvent such as polyethylene glycol or a salt such as ammonium sulfate. Other suitable separation techniques include, without limitation, the fluorescein antibody magnetizable particle method described in Rattle, et al. (1984) <u>Clin. Chem.</u> 30:1457-1461, and the double antibody magnetic particle separation as described in U.S. Pat. No. 4,659,678.

Methods for linking proteins or their fragments to the various labels have been extensively reported in the literature and do not require detailed discussion here. Many of the techniques involve the use of activated carboxyl groups either through the use of carbodiimide or active esters to form peptide bonds, the formation of thioethers by reaction of a mercapto group with an activated halogen such as chloroacetyl, or an activated olefin such as maleimide, for linkage, or the like. Fusion proteins will also find use in these applications.

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Another diagnostic aspect of this invention involves use of oligonucleotide or polynucleotide sequences taken 20 from the sequence of a chemokine. These sequences can be used as probes for detecting levels of the ligand message in samples from patients suspected of having an abnormal condition, e.g., an inflammatory or developmental problem. The preparation of both RNA and DNA nucleotide sequences, the labeling of the sequences, and the preferred size of the sequences has received ample description and discussion in the literature. Normally an oligonucleotide probe should have at least about 14 nucleotides, usually at least about 18 nucleotides, and the polynucleotide probes may be up to several kilobases. 30 Various labels may be employed, most commonly radionuclides, particularly 32P. However, other techniques may also be employed, such as using biotin modified nucleotides for introduction into a 35 polynucleotide. The biotin then serves as the site for binding to avidin or antibodies, which may be labeled with a wide variety of labels, such as radionuclides, fluorescers, enzymes, or the like. Alternatively, antibodies may be employed which can recognize specific

duplexes, including DNA duplexes, RNA duplexes, DNA-RNA hybrid duplexes, or DNA-protein duplexes. The antibodies in turn may be labeled and the assay carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected. The use of probes to the novel anti-sense RNA may be carried out in any conventional techniques such as nucleic acid hybridization, plus and minus screening, recombinational probing, hybrid released translation (HRT), and hybrid arrested translation (HART). This also includes amplification techniques such as polymerase chain reaction (PCR).

Diagnostic kits which also test for the qualitative or quantitative presence of other markers are also contemplated. Diagnosis or prognosis may depend on the combination of multiple indications used as markers. Thus, kits may test for combinations of markers. See, e.g., Viallet, et al. (1989) Progress in Growth Factor Res. 1:89-97.

X. Receptor

Having isolated a ligand binding partner of a specific interaction, methods exist for isolating the - 25 counter-partner. See, Gearing, et al EMBO J. 8:3667-4676 or McMahan, et al. (1991) <u>FMBO J.</u> 10:2821-2832. For example, means to label a chemokine without interfering with the binding to its receptor can be determined. For example, an affinity label can be fused to either the 30 amino- or carboxy-terminus of the ligand. An expression library can be screened for specific binding of chemokine, e.g., by cell sorting, or other screening to detect subpopulations which express such a binding component. See, e.g., Ho, et al. (1993) Proc. Nat'l Acad. Sci. 90:11267-11271. Alternatively, a panning 35 method may be used. See, e.g., Seed and Aruffo (1987) Proc. Nat'l. Acad. Sci. 84:3365-3369.

Protein cross-linking techniques with label can be applied to isolate binding partners of a chemokine. This

would allow identification of protein which specifically interacts with a chemokine, e.g., in a ligand-receptor like manner.

In various embodiments, new receptors designated DC CR and M/DC CR were isolated. The sequences of the human constructs are set forth in SEQ ID NOS: 10 and 12. Similar means for making variants and fragments, at the nucleotide level or at the protein level, and making antibodies will be available as described above, directed primarily to the chemokine embodiments. Many similar or related uses to the ligands will be applied to the receptors, as specific binding reagents. In particular, methods will be applied to screening for specific ligands for each receptor. Many uses, including kits, will also be available through analogous techniques.

The broad scope of this invention is best understood with reference to the following examples, which are not intended to limit the invention to specific embodiments.

20 EXAMPLES

I. General Methods

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Some of the standard methods are described or referenced, e.g., in Maniatis, et al. (1982) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor 25 Laboratory, Cold Spring Harbor Press; Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual, (2d ed.), vols 1-3, CSH Press, NY; Ausubel, et al., Biology, Greene Publishing Associates, Brooklyn, NY; or Ausubel, et al. (1987 and Supplements) Current Protocols in Molecular 30 Biology, Greene/Wiley, New York; Innis, et al. (eds.) (1990) PCR Protocols: A Guide to Methods and Applications Academic Press, N.Y. Methods for protein purification include such methods as ammonium sulfate precipitation, column chromatography, electrophoresis, 35 centrifugation, crystallization, and others. See, e.g., Ausubel, et al. (1987 and periodic supplements); Deutscher (1990) "Guide to Protein Purification" in Methods in Enzymology, vol. 182, and other volumes in

this series; and manufacturer's literature on use of protein purification products, e.g., Pharmacia, Piscataway, N.J., or Bio-Rad, Richmond, CA. Combination with recombinant techniques allows fusion to appropriate segments, e.g., to a FLAG sequence or an equivalent which can be fused via a protease-removable sequence. See, e.g., Hochuli (1989) Chemische Industrie 12:69-70; Hochuli (1990) "Purification of Recombinant Proteins with Metal Chelate Absorbent" in Setlow (ed.) Genetic

Engineering. Principle and Methods 12:87-98, Plenum Press, N.Y.; and Crowe, et al. (1992) Olaexpress: The High Level Expression & Protein Purification System QUIAGEN, Inc., Chatsworth, CA.

FACS analyses are described in Melamed, et al.

(1990) Flow Cytometry and Sorting Wiley-Liss, Inc., New
York, NY; Shapiro (1988) Practical Flow Cytometry Liss,
New York, NY; and Robinson, et al. (1993) Handbook of
Flow Cytometry Methods Wiley-Liss, New York, NY.

20 II. Isolation and characterization of chemokine cDNAs A. TECK

The TECK was isolated from a cDNA library made from thymus cells from a RAG-1 "knockout" mouse. See, Mombaerts, et al. (1992) Cell 68:869-877. Individual

- cDNA clones were sequenced using standard methods, e.g., the Taq DyeDeoxy Terminator Cycle Sequencing kit (Applied Biosystems, Foster City, CA), and the TECK sequence was identified and further characterized. Computer analyses with other C-C chemokine family members revealed
- significant homology at the amino acid levels with other chemokines. The nucleotide sequence for mouse is provided in SEQ ID NO: 1, encoding a polypeptide of about 144 amino acids. The signal sequence should run from 1 (met) to about 23 (ala), and removal of the signal
- sequence should provide one natural mature sequence beginning at 24 (gln). Additional processing may occur in a physiological system.

The sequence is notable in having a longer carboxy-terminal tail than most other CC chemokines. TECK

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exhibits one glycosylation site, and several AAMP, PKC, and CK2 phosphorylation sites.

B. $MIP-3\alpha$

The MTP-3α was isolated from a cDNA library made from human monocytes activated with LPS and IFN-γ
Individual cDNA clones were sequenced using standard methods, and the MIP-3α sequence was identified and further characterized. The nucleotide sequence is set forth in SEQ ID NO: 5, encoding a polypeptide of at least about 89 amino acids. The signal sequence should run from about 1 (met) to 21 (cys), and removal of the signal sequence should provide one natural sequence beginning with gly. Additional processing may occur in a physiological system.

C. MIP-3B

The MIP-3 β was isolated from a cDNA library made from human fetal lung cells. Individual cDNA clones are sequenced using standard methods, and the MIP-3 α sequence was identified and further characterized. The nucleotide sequence is set forth in SEQ ID NO: 7, encoding a polypeptide of about 98 amino acids. The signal sequence should run from about 1 (met) to about 21 (ser), and removal of the signal sequence should provide one mature natural sequence beginning from gly. Additional processing may occur in a physiological system.

D. Dendritic Cell Receptor for chemokine; DC CR
The DC CR was isolated from RNA made from dendritic
cells isolated from CD34+ cord blood cells, isolated by
standard procedure. It was also isolated from
eosinophils using degenerate PCR primers of the TM2 and
TM7 segments, which are often conserved among chemokine
receptors. These eosinophils were isolated by taking
PBLs, depletion of red blood cells by lysis, and negative
selection of CD16 to remove neutrophils.

Sequencing of the PCR fragments indicated a potential novel receptor, and the fragment was used to isolate a full length clone by hybridization. Clone isolates were sequenced using standard methods, and the DC CR sequence was identified and further characterized.

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The nucleotide sequence is provided in SEQ ID NO: 9, encoding a polypeptide of about 365 amino acids. The transmembrane segments, determined by homology to the IL-8 B receptor, are about: TM1 from 39 (leu) to 64 (phe): TM2 from 76 (leu) to 96 (ser); TM3 from 111 (leu) to 132 (met); TM4 from 151 (thr) to 176 (phe); TM5 from 207 (gly) to 229 (val); TM6 from 246 (val) to 270 (ala); and TM7 from 291 (val) to 319 (leu). The amino terminal segment is probably an extracellular segment, and the 10 others would be between TM2 and TM3; and TM4 and TM5; and TM6 and TM7. The intracellular segments should then run between TM1 and TM2; TM3 and TM4, TM5 and TM6, and the carboxy terminus from the end of TM7. Additional processing may occur in a physiological system.

The implication of chemokine receptors in retroviral infection suggests that the receptor may be critical for infection. Antibodies which block infection may be routinely screened, and developed for therapeutic uses.

E. Monocyte/Dendritic Cell Receptor for chemokine; 20 M/DC CR

The M/DC CR was isolated from a cDNA library made from human monocyte cells cultured for 2.5 to t h in medium containing IFN- γ (10 ng/ml), LPS (1 μ g/ml), anti-IL-4 monoclonal antibody (5 μ g/ml), and anti-IL-10 monoclonal antibody (5 μ g/ml). Individual cDNA clones were sequenced using standard methods, and the M/DC CR sequence was identified and further characterized. nucleotide sequence is set forth in SEQ ID NO: 11, encoding a polypeptide of about 356 amino acids. The 30 transmembrane segments, should be about as follows: TM1 from 52 (leu) to 76 (val); TM2 from 86 (asn) to 107 (ala); TM3 from 117 (ile) to 138 (val); TM4 from 157 (val) to 182 (tyr); TM5 from 211 (phe) to 233 (val); TM6 from 251 (leu) to 275 (phe); and TM7 from 296 (ile) to 35 315 (leu). As for the DC CR, the amino terminal segment is probably an extracellular segment, and the others would be between TM2 and TM3; and TM4 and TM5; and TM6 and TM7. The intracellular segments should then run

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between TM1 and TM2; TM3 and TM4, TM5 and TM6, and the carboxy terminus from the end of TM7.

III. Preparation of antibodies

Many standard methods are available for preparation of antibodies. For example, synthetic peptides may be prepared to be used as antigen, administered to an appropriate animal, and either polyclonal or monoclonal antibodies prepared. Short peptides, e.g., less than about 10 amino acids may be repeated, while longer 10 peptides may be used alone or conjugated to a carrier. For example, with the M/DC CR, animals were immunized with peptides corresponding to amino acid sequences from 18-44 (starting with LAP and ending with KYD; a fragment towards the amino terminus) and from 183-204 (starting with KPQ and ending with PAD; corresponding to an extracellular loop), see SEQ ID NO: 13. Highest specificity will result when the polypeptides are selected from portions which are most unique, e.g., not form conserved sequence regions. The animals may be used to collect antiserum, or may be used to generate monoclonal antibodies.

Antiserum was determined useful for ELISA, and will be evaluated for utility as immunoprecipitation or Western blot analysis. Monoclonal antibodies will also be evaluated for those same uses.

The antibodies provided will be useful as immunoaffinity reagents, as detection reagents, for immunohistochemistry, and as therapeutic reagents.

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IV. Assays for chemotactic activity of chemokines. Chemokine proteins are produced, e.g., in COS cells transfected with a plasmid carrying the chemokine cDNA by electroporation. See, Hara, et al. (1992) EMBO J. 35 10:1875-1884. Physical analytical methods may be applied, e.g., CD analysis, to compare tertiary structure to other chemokines to evaluate whether the protein has likely folded into an active conformation. After transfection, a culture supernatant is collected and

subjected to bioassays. A mock control, e.g., a plasmid carrying the luciferase cDNA, is used. See, de Wet, et al. (1987) Mol. Cell. Biol. 7:725-757. A positive control, e.g., recombinant murine MIP-1 α from R&D Systems (Minneapolis, MN), is typically used. Likewise, antibodies may be used to block the biological activities, e.g., as a control.

Lymphocyte migration assays are performed as previously described, e.g., in Bacon, et al. (1988) Br.

10 J. Pharmacol. 95:966-974. Murine Th2 T cell clones, CDC-25 (see Tony, et al. (1985) J. Exp. Med. 161:223-241) and HDK-1 (see Cherwinski, et al. (1987) J. Exp. Med. 166:1229-1244), made available from R. Coffman and A. O'Garra (DNAX, Palo Alto, CA), respectively, are used as controls.

Ca2+ flux upon chemokine stimulation is measured according to the published procedure described in Bacon, et al. (1995) <u>J. Immunol</u>. 154:3654-3666.

Maximal numbers of migrating cells in response to 20 MIP-1α typically occur at a concentration of 10-8 M, in agreement with original reports for CD4+ populations of human T cells. See Schall (1993) J. Exp. Med. 177:1821-1826. A dose-response curve is determined, preferably giving a characteristic bell shaped doseresponse curve.

After stimulation with C-C chemokines, lymphocytes generally show a measurable intracellular Ca2+ flux. MIP-1 α is capable of inducing immediate transients of calcium mobilization. Typically, the levels of chemokine used in these assays will be comparable to those used for the chemotaxis assays (1/1000 dilution of conditioned supernatants).

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Retroviral infection assays have also been described, and recent description of certain chemokine receptors in retroviral infection processes may indicate that similar roles may apply to the DC CR and/or M/DC CR. See, e.g., Balter (1996) <u>Science</u> 272:1740 (describing evidence for chemokine receptors as coreceptors for HIV); and Deng, et al. (1996) <u>Nature</u> 381:661-666.

Expression analysis of chemokine/receptor genes ٧. RNA blot and hybridization are performed according to the standard method in Maniatis, et al. (1982) Molecular Cloning: A Laboratory Manual Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY. An appropriate fragment of a cDNA fragment is selected for use as a probe. To verify the amount of RNA loaded in each lane, the substrate membrane is reprobed with a control cDNA, e.g., glyceraldehyde 3-phosphate dehydrogenase (G3PDH) cDNA (Clontech, Palo Alto CA). Analysis of mRNA from the appropriate cell source using the probe will determine the natural size of message. It will also indicate whether different sized messages exist. The messages will be subject to analysis after isolation, e.g., by PCR or hybridization techniques. Northern blot analysis may be performed on many different mRNA sources. However, in certain cases, cDNA libraries may be used to evaluate sources which are difficult to prepare. A "reverse Northern" uses cDNA inserts removed from vector, but multiplicity of bands may reflect either different sized messages, or may be artifact due to incomplete reverse transcription in the preparation of the cDNA library. In such instances, verification may be appropriate by standard Northern analysis. Similarly, Southern blots may be used to evaluate species distribution of a gene. The stringency of washes of the blot will also provide information as to the extent of homology of various species counterparts.

Tissue distribution, and cell distribution, may be evaluated by immunohistochemistry using antibodies. Alternatively, in situ nucleic acid hybridization may also be used in such analysis.

A TECK

The TECK was isolated from a RAG-1 "knockout" mouse. This animal is characterized by a great predominance of pro-T or pre-T cells, lacking more mature T cells after the block point of T cell receptor rearrangement. This suggests a role in very early T cell development, likely expressed by pro-T or pre-T cells, thymic stromal cells, and possibly macrophages, epithelial, and dendritic cells. This comports with the observation that tissue distribution

Table 2: mTECK mRNA expression in tissues and cells

cDNA libraries		northern blot	
cell type or tissue	neg pos	cell type or tissue	neg pos
Th2 CD4+ T cells	х	heart	х
Th1 CD4+ T cells	X	brain	x
Lung	X	spleen	X
L cells	x	lung	X
RAG-1 KO lung	X	liver	X
RAG-1 KO heart	X	skeletal muscle	x
RAG-1 KO brain	X (+)	kidney	x
RAG-1 KO spleen	X	testis	x ·
RAG-1 KO kidney	· X	thymus	X (+++)
RAG-1 KO testis	X (+)	small intestine	X (++)
RAG-1 KO thymus	X (+++)		Х (++)
RAG-1 KO liver	X (+)	CD4-8+ thymocytes R/A	X
CD4-8- thymocytes	X	CD4-8- thymocytes R/A	X.
A20-J B-cell lymphoma	x	B220+ splenocytes R/A	X
BW CD4-8-3- hybridoma	X	Thy-1+ splenocytes R/A	X
pro-T cells	X (+)	1G18LA macrophages R/A	X
pre-T cells	x	primary thymic stroma R/A	X
30-R bone marrow stroma	X	3D.1 thymic epithelial R/A	X
D10 T-cell hybridoma	X	MTSC-C thymic epithelial	X
CTLL T-cell clone	X	30.R bone marrow stroma	X
peritoneal macrophages	Х		46
splenic dendritic cells	X		4

Analysis of mTECK mRNA was carried out as described. + to +++ indicates the relative intensity of the signal. R/A: resting or activated.

Species analysis indicated positive signals by hybridization in human, rat, and hamster DNA. Tissue distribution analysis suggests that the gene is expressed in human small intestine, which also is a tissue which supports T cell differentiation.

The combination of the structure and distribution of this chemokine suggests a role in T cell development, which normally occurs in the thymus.

B. $MIP-3\alpha$

- The MIP-3 α was identified from a cDNA library made from human monocytes activated with LPS and IFN- γ , in the presence of anti-IL-10. It has also been detected in pancreatic islet cells, fetal lung, and hepatic HEPG2 cells.
- The gene is expressed in HL-60 (promyelocytic leukemia); S3 (HeLa cell); K562 (chronic myelogenous leukemia); MOLT-4 (lymphblastic leukemia); Raji

(Burkitt's lymphoma); SW480 (colorectal adenocarcinoma); A549 (lung carcinoma); and G361 (melanoma) cell lines, as determined by probing on a tissue blot from CLONTECH. Tissue expression gave a positive signal in lymph node, appendix, peripheral blood lymphocytes, fetal liver, and fetal lung; but no detectable signal in spleen, bone marrow, brain, and kidney.

The main transcript appears to be about 1.2 kb, with two additional transcript sizes in fetal lung RNA. Among the various tissues, transcript sizes of 1.8, 2.7, and 4.2 kb were detected.

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Positive signals were also detected in the following cDNA libraries: dendritic cells activated with LPS, but not when activated with GM-CSF and IL-4; monocytes treated with LPS, IFN-γ, and anti-IL-10, but not when treated with LPS, IFN-γ, and IL-10; and activated PBMC.

These expression data implicate this chemokine in inflammatory responses upon cell activation. The lymph nodes, appendix, and PBL are sites where inflammatory processes take place. The MIP-3 α may exert its effects on monocytes and cells involved in inflammatory events. Other structural features implicate this chemokine in eosinophil and lung physiology, e.g., asthma indications. Thus, an antagonist of the chemokine, e.g., an antibody, may be important for treatment of asthmatic conditions. Also, IL-10 appears to inhibit MIP-3 α expression.

The human MIP-3 α is a ligand for the DC CR. Thus, a positive control exists for the Ca++ flux assay for that receptor. This allows for the further screening of agonist ligands for the DC CR. Moreover, the DC CR was isolated from eosinophil cDNA, and observations have been made that eosinophils migrate to MIP-3 α in vitro. This suggests that the MIP-3 α interaction with the DC CR is important in recruitment of eosinophils, as occurs with the eotaxin ligand and the CCR3. As such, antagonists of the MIP-3 α interaction with the DC CR will likely be useful in inhibiting eosinophilia, particularly in the lung, or lung inflammation. These may accompany asthmatic or other pulmonary conditions.

Antagonists to MIP-3α may be made either with antibodies, or other binding compositions which inhibit receptor interaction. The antibodies may be to the ligand, MIP-3α itself, or to the binding portions of the receptor, DC CR. Muteins of the chemokine may block receptor interaction, and with a positive control, chemokine muteins may be screened for variations which compete with the wild type chemokine at various concentrations. See, e.g., Kenakin (1987)

Pharmacological Analysis of Drug-Receptor Interaction
Raven Press, NY; Arunlakshana and Schild (1959) Br. J.
Pharmacol. 14:48-58; Black (1989) Science 245:486-493;
Zurawski, et al. (1986) J. Immunol. 137:3354-3360;
Zurawski and Zurawski (1988) EMBO J. 7:1061-1069;
Zurawski and Zurawski (1992) EMBO J. 11:3905-3910; Imler and Zurawski (1992) J. Biol. Chem. 267:13185-13190.

C. MIP-3B

The MIP-3 β was identified in a cDNA library made from human monocytes activated with LPS and IFN- γ , in the presence of anti-IL-10. Its distribution in other cells and tissues has not been fully determined.

- D. Dendritic Cell Receptor for chemokine; DC CR
 The DC CR was isolated from a cDNA library made from
 a dendritic cell cDNA library. It appears to be
 expressed in certain T cells, spleen cell subsets, NK
 cells, and other cell populations enriched in dendritic
 cells, including CD1a+, CD14+, and CD1Aa+ cells. It did
 not give a detectable signal in TF1, Jurkat, MRC5, JY, or
 U937 cells.
- Being found on dendritic cells, its ligand, including the MIP-3α, may be important in attracting appropriate cells for the initiation of an immune response. MIP-3α is therefore likely to attract dendritic cells, leading to initiation of the immune response. Pulmonary physiology is suggested, both from the distribution of the receptor and ligand. The receptor may be also present in other cells important in such responses.

E. Monocyte/Dendritic Cell Receptor for chemokine;
M/DC CR

The M/DC CR was isolated from a cDNA library made from primary monocyte cells activated with LPS and IFN- γ but subtracted with known high abundance genes from those cells. The abundance of this gene is probably less than about 1% of message from those cells.

Tissue expression gave a positive signal in spleen, PBL, lung, placenta, and small intestine; but no detectable signal in brain, liver, kidney, and muscle. This distribution suggests a hematopoietic role.

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There appears to be one main transcript, but the existence of additional or alternatively spliced messages has not been eliminated.

Positive signals were also detected in the following cDNA libraries: monocytes and dendritic cells; but signals were not detectable in CD8+ T cells, or in either resting or activated splenocytes, gamma-delta T cells, NK cells, or B cells. Immunohistochemistry will be performed to confirm absence in the T cell and B cell compartments and to check in tonsil, particularly in view of location in spleen and placenta. The relatively restricted distribution on monocytes and dendritic cells leads both to its designation, and suggests a functional role in those cell types, which are important in the initiation of immune responses through their ability to process and present antigen to T cells.

VI. Specific Characterization of TECK

A novel CC chemokine was identified in the thymus of mouse and human and was designated TECK as Thymus Expressed ChemoKine. TECK has weak homology with other CC chemokines and maps to mouse chromosome 8. Besides the thymus, mRNA encoding TECK was detected at substantial levels in the small intestine and at low levels in the liver. The source of TECK in the thymus was determined to be thymic dendritic cells, while in contrast bone marrow-derived dendritic cells do not express TECK. The murine TECK recombinant protein showed

chemotactic activity for activated macrophages, dendritic cells and thymocytes. We conclude that TECK represents a novel thymic dendritic cell-specific CC chemokine which is possibly involved in T-cell development.

Chemokines belong to a family of small peptides (6-5 15 kDa) whose best described biological function is to control the migration of certain leukocyte populations to localized sites of inflammation. Baggiolini, et al. (1994) Adv. in Immun. 55:97-179; Schall and Bacon (1994) 10 Curr Opin Immun 6:865-873; Hedrick and Zlotnik (1996) Curr. Opin. Immunol. 8:343-347. In the last few years many new members of the chemokine super family have seen the characterized. Initially, new chemokines were identified through their chemotactic effects on leukocytes (Baggiolini et al. (1994); Schall and Bacon 15 (1994)) and were isolated mainly from blood leukocytes or cell lines. More recently, approaches based on the selective cloning of secreted molecules by signal sequence trap (Tashiro, et al. (1993) Science 261:600-20 603; Imai, et al. (1996) <u>J. Biol. Chem.</u> 271:21514-21521) or on the exploitation of public and private databases of expressed sequence tags (EST) through bioinformatics (Hieshima, et al. (1997) <u>J. Biol. Chem.</u> 272:5846~5853; Patel, et al. (1997) J. Exp. Med. 185:1163-1172; and 25 Rossi, et al. (1997) J. Immunol. 158:1033-1036), have allowed the rapid identification of novel chemokines based on sequence and structural homologies. These approaches take advantage of the fact that most of the chemokines are secreted factors whose protein sequence 30 contain four conserved cysteines (Schall (1994) *The Chemokines" pp. 419-460 in Thomson (eds.) The Cytokine Handbook, Academic Press, New York. The CXC or a chemokine family has the two first amino-terminal cysteines separated by a non-conserved amino acid. In 35 the CC or β chemokine family, these two cysteines are consecutive. A third type of chemokine, the C or γ family, is represented by lymphotactin, which conserves

two cysteines (1 and 3) instead of the original four (Kelner, et al. (1994) <u>Science</u> 266:1395-1399). Finally,

a recently identified chemokine with three amino acids separating the first two cysteines defines a fourth CX3C family (Bazan, et al. (1997) Nature 385:640-644).

Interestingly, some of the new chemokines discovered show a relatively restricted pattern of expression (Imai et al. (1996); Hieshima et al. (1997)). These new approaches may lead to the discovery of tissue- or cellspecific chemokines. In addition, new biological evidence exists for important new roles of chemokines in hemopoiesis (Cook (1996) J. Leukoc. Biol. 59:61-66; and Nagasawa, et al. (1996) Nature 382:635-638) and the control of viral infections including HIV (Cocchi, et al. (1995) Science 270:1811-1815; and Cook, et al. (1995) Science 269:1583-1585). Thus, the molecular cloning of novel chemokines through DNA-based strategies may uncover novel proteins belonging to the chemokine super family but whose physiological role goes beyond the control of inflammation.

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In an attempt to identify novel genes involved in Tcell development, we analyzed a cDNA library from the thymus of Recombinase Activation Gene-1 (RAG-1) deficient mice. We identified a novel CC chemokine designated TECK for Thymus Expressed ChemoKine, based on sequence homology with other known chemokines. We subsequently isolated the human homologue of TECK. The pattern of expression of TECK mRNA is highly restricted to the thymus and small intestine in both human and mouse. Moreover, in the mouse thymus, TECK protein is produced by dendritic cells while splenic dendritic cells do not express TECK mRNA. Recombinant TECK showed chemotactic 30 activity on thymocytes, macrophages, THP-1 cells and dendritic cells, while it was inactive on peripheral lymphocytes and neutrophils. The restricted pattern of expression of TECK together with its biological 35 properties suggests a role for this novel dendritic cellspecific chemokine in T-cell development.

A. Cloning and structural analysis of mouse TECK A directional cDNA library was made from RAG-1 deficient mouse thymus and analyzed by random sequencing.

One of the clones contained an open reading frame with significant homology to previously described CC chemokines. The full-length cDNA contains 1037 bp including an open reading frame of 426 bp encoding a protein of 142 amino acids and will be identified in this report as mTECK (see SEQ ID NO: 1). In the 3' untranslated region, there is one unique polyadenylation signal consistent with the single mRNA species observed in northern blots. The mTECK cDNA does not contain any ATTTA transcript destabilization motif (Shaw and Kamrn 10 (1986) Cell 46:659-667). The comparison of the amino acid sequence of mTECK with previously described murine CC chemokines demonstrates the conservation of the four cysteines present in all these chemokines. However, mTECK shows few additional identities with these 15 proteins.

B. Cloning and molecular characterization of human TECK To investigate the possible existence of a gene homologous to mTECK in other mammalian species, a Southern blot with genomic DNA from various species was 20 hybridized with the mTECK cDNA probe. Under high stringency conditions, hybridizing bands were detected in mouse, rat, hamster and human genomic DNAs. Interestingly, a single band was detected in human, 25 suggesting that a single gene encodes for TECK in this species. The multiple bands present in mouse, rat and hamster could be the result of an internal EcoRI site within the TECK gene. Alternatively, the TECK gene may have been duplicated in these species. 30

In order to clone the human homologue of mTECK, a blot of cDNAs from a panel of human cDNA libraries was hybridized with the mTECK cDNA probe. A signal was observed in a fetal small intestine cDNA library. Screening of this library with the mTECK probe allowed the isolation of several identical clones of 1012 bp with an open reading frame of 453 bp encoding a protein of 151 amino acids. This protein had a much higher degree of homology at the nucleic acid level (71% nucleic acid identity for the open reading frame and 49.3% amino acid

identity) to mTECK than to other known CC chemokines and was thus designated as hTECK.

C. Chromosomal location of mTECK

It has been shown that the genes encoding for most 5 chemokines are clustered in the genome. The genes encoding CC chemokines cluster on mouse chromosome 11 and Schall (1994); and Hedrick human chromosome 17q11-12. and Zlotnik (1996). The chromosomal location of mTECK (designated gene symbol Teck) was determined by inter specific backcross analysis between ([C57B1/6J X M. 10 spretus]F1 X C57B1/6J) mice. Jenkins, et al. (1982) J. Virol. 43:26-36. The mapping results indicated that the Teck locus is located on the proximal part of mouse chromosome 8. Although the chromosomal location of the human Teck locus could not be determined, this region of 15 mouse chromosome 8 is syntenic to the human 19p13.3 and 13q34 regions. However, the Teck locus is also very close to a region syntenic to human chromosome 4. The closest known gene, Insr, encodes the insulin receptor and the genetic distance between Teck and Insr was 20 estimated at 0.9 \pm 0.9 cM. We have compared our inter specific map of chromosome 8 with a composite mouse linkage map that reports the map location of many uncloned mouse mutations (Mouse Genome Database, The Jackson Laboratory, Bar Harbor, ME). Teck resides in a 25 region of the composite map that lacks mouse mutations with a phenotype that might be expected for an alteration in this locus.

D. Analysis of mTECK and hTECK mRNA distribution in cells and tissues

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An analysis of the distribution of mTECK mRNA in tissues and cells by northern blotting or by Southern blotting of mouse cDNA libraries revealed that mTECK was expressed at significant levels only in the thymus and to a lesser extent in small intestine (Table 2). Weak expression of mTECK mRNA was observed in brain, testis, and liver RAG-1-/- cDNA libraries. Interestingly, mTECK mRNA was detected in a cDNA library of activated pro-T cells. Pro-T cells represent an early stage of intra-

thymic T-cell progenitors, not fully committed to the T-cell lineage since they can give rise to NK and dendritic cells. Moore and Zlotnik (1995) <u>Blood</u> 86:1850-1860; Wu, et al. (1996). <u>J. Exp. Med.</u> 184:903-911). In contrast, mTECK mRNA was undetectable in resting or activated thymocytes, peripheral T or B cells, macrophages, PBLs, splenic dendritic cells and in all other tissues tested, with the exception of spleens recovered from mice injected with LPS (Table 2). Interestingly, mTECK mRNA was detected by PCR in fetal thymuses of day 14 of gestation, indicating that mTECK is expressed in the

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thymus at the earliest stages of T-cell development. The distribution of hTECK mRNA was similarly analyzed. As with mouse, hTECK mRNA expression was 15 highly restricted to the thymus and small intestine. Weak expression was also detected in inflamed tonsil and fetal spleen, but at much lower levels than that observed in the thymus since this particular blot was exposed for a long time. Importantly, hTECK mRNA was absent from a 20 series of cDNA libraries from dendritic cells derived in vitro from bone marrow CD34+ progenitors cells or peripheral blood monocytes. In addition, hTECK mRNA was also absent from libraries of monocyte-derived dendritic cells stimulated with LPS or a combination of TNF- α , IL- 1α and monocyte supernatant for 4 and 16 hours. Collectively, these data indicate that TECK mRNA is. specifically expressed at high levels in thymus and small intestine in vivo.

The abundance of mTECK mRNA expression in RAG-1
deficient thymus and its absence in thymic T cells
suggested that mTECK was expressed by a thymic stromal
component in normal mice. We performed in situ mRNA
hybridization with sense or antisense mTECK probes

generated by PCR. Thymic sections hybridized with the
sense probe (negative control) demonstrated no specific
staining while sections hybridized with the anti-sense
probe at the same concentration showed specific staining
in the thymic medulla. At higher magnification, positive

cells appeared to have a non-lymphoid morphology with processes surrounding lymphoid cells. This experiment indicated that, in vivo, mTECK mRNA is expressed by a non-lymphoid component of the medullary stroma, possibly dendritic cells.

The thymic stroma is mainly composed of epithelial cells, macrophages, dendritic cells and fibroblasts, together with a network of vascular and nervous tissue. Boyd, et al. (1993) Immunol. Today 14:445-459. Since we previously failed to detect mTECK mRNA expression in thymic epithelial or macrophage cell lines with our without activation with IFN-\gamma (Table 2), we sorted thymic dendritic cells based on their high expression of MHC class II and CD11c (N-418 antibody). Analysis of mTECK expression by RT-PCR revealed that freshly isolated MHC class II+ CD11c+ thymic dendritic cells expressed mTECK mRNA while the MHC class II+ CD11c- subset was negative. In contrast, mTECK mRNA was undetectable in a cDNA library made from freshly isolated splenic dendritic cells (Table 2).

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We then performed immunostaining of thymic sections and purified thymic dendritic cells with a polyclonal antibody raised against a decapeptide corresponding to the C-terminus of mTECK. This polyclonal antibody reacted with recombinant mTECK in ELISA and western blot while normal rabbit serum was negative. In thymic sections, the polyclonal anti-peptide antibody reacted with a stromal component of the thymic medulla consistent with the in situ hybridization data while staining with normal rabbit serum was negative. Interestingly, the antibody also reacted weakly with some endothelial cells, raising the possibility that mTECK can be produced by the thymic endothelium. Finally, the anti-mTECK polyclonal antibody stained sorted thymic dendritic cells, while the control serum was negative. High magnification clearly showed intra-cellular staining of cells with characteristic dendritic morphology. Taken together, these results indicate that thymic dendritic cells and

possibly thymic endothelial cells are producing TECK in vivo.

F. Chemotactic activities of mTECK protein

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To evaluate the biologic properties of mTECK, a recombinant protein with an N-terminal FLAG peptide was obtained in a bacterial expression system. In some experiments, a recombinant mTECK protein with a C-terminal FLAG was used and similar results were obtained. Interestingly, mTECK induced the migration of mouse

thymocytes (Figure 1A). The optimal response was obtained with a dose of 10 ng/ml TECK. Cell migration was determined to be chemotaxis and not chemokinesis through the checkerboard analysis. Furthermore, it is established that chemokines bind to specific receptors

that are coupled through heterotrimeric G proteins to intra-cellular signal-transducing pathways. Murphy (1994) Annu. Rev. Immunol. 12:593-633. To determine whether the chemotaxis of thymocytes involved a G protein-coupled receptor, cells were incubated prior to

20 the assay with 10 ng/ml pertussis toxin which ADP-ribosylates Gαi-proteins. Katz, et al. (1992) Nature 360:686-689. This pre-treatment completely abrogated the chemotactic response of thymocytes to mTECK (Figure 1A).

The recombinant mTECK protein also induced the 25 migration of human monocytic THP-1 cells activated for 16 hours with IFN-γ (Figure 1B), while it was not significantly active on resting THP-1 cells. This experiment showed that mTECK is active on human cells. In addition, mTECK induced activated mouse peritoneal 30 macrophages to migrate as well as highly purified mouse splenic dendritic cells (Figure 1B). In all these experiments, the optimal dose of mTECK was 10 ng/ml. contrast, no chemotaxis was observed with bone marrow cells, purified neutrophils, splenic B cells, splenic T cells or IL-2 activated RAG-1 deficient mouse splenocytes lacking mature T and B lymphocytes (Mombaerts, et al. (1992) Cell 68:869-877) and therefore enriched in NK cells. These data are consistent with the absence of in vivo accumulation of neutrophils, monocytes or

lymphocytes 2 and 5 h following an intra-peritoneal injection of 10 µg mTECK. Collectively, these data indicate that TECK is a chemotactic factor for

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thymocytes, macrophages and dendritic cells. TECK, a distant member of the CC chemokine family In this report, we describe the molecular isolation and characterization of TECK, a novel mouse and human CC chemokine. Analysis of its predicted amino acid sequence showed that TECK is distantly related to previously described CC chemokines. Conservation of particular amino acids among most CC chemokines may be related to their functional importance. Beall, et al. (1992) J. Biol. Chem. 267:3455-3459; and Lusti-Narasimhan, et al. (1995) J. Biol. Chem. 270:2716-2721. In particular, a tyrosine residue between the second and third cysteines has been shown to be critical for monocyte chemotaxis (in position 50) (Beall et al. (1992)). While TECK does not have a tyrosine at this particular position, it has one in position 52 that may have the same function, since TECK is chemotactic for activated monocytes. In addition to these differences in the primary structure, the gene encoding TECK maps on chromosome 8 in the mouse, unlike most other CC chemokines which are clustered on chromosome 11. This is not the first report of an unusual chromosomal location for a CC chemokine. We have cloned the human CC chemokine MIP-3B and showed that its gene was on chromosome 9 rather than 17 (Rossi, et al. (1997)), and the gene encoding the novel human CC chemokine MIP-3 α /LARC (Rossi, et al. (1997)) has been mapped on chromosome 2 (Hieshima, et al. (1997)). It is likely that the CC chemokines on chromosome 11 in the mouse and 17 in human have been generated through gene duplication of a primordial chemokine. Our results suggest that TECK may have been generated at an earlier stage during evolution. In this regard, the TECK gene 35 may have evolved to ensure functions similar to other CC chemokines with a distant primary structure but through similar receptor(s) as dictated by its secondary and

tertiary structures. Alternatively, the receptor(s) and

physiological role of TECK may be unique among chemokines.

H. TECK expression and function is associated with T-cell development

5 We observed that TECK was strongly expressed in the thymus which is the primary lymphoid organ where T-cell development takes place. Recently, another CC chemokine highly expressed in the thymus, TARC, has been identified. Imai, et al. (1996). However, TARC is also 10 expressed in lung and colon as well as activated PRMC (Imai, et al. (1996)) while TECK was absent from these tissues. Besides the thymus, numerous reports indicate that T cell development can occur in the small intestine (Poussier and Julius (1994) Annu. Rev. Immunol. 12:521-15 553) where TECK is also expressed. Interestingly, the liver has also been suggested to support T-cell development to some extent (Abo, et al. (1994) Int. Rev. Immunol. 11:61-102) and we observed a low TECK expression

Immunol. 11:61-102) and we observed a low TECK expression
in a liver cDNA library. These data show that TECK
20 expression correlates with organs that support T-cell
development.

While many molecular and cellular aspects of T-cell differentiation are well documented, the precise role of chemokines in T-cell development is still unknown.

Recently, it has been shown that the bone marrow stromaderived CXC chemokine SDF-1 is important for B lymphopoiesis and myelopoiesis since SDF-1 -/- mice are impaired for these functions (Nagasawa, et al. (1996)). Similarly, it is likely that chemokines act at different steps of T-cell differentiation. Chemokines, together with the expression of appropriate adhesion molecules, may dictate the migration of uncommitted progenitors from the bone marrow to other anatomic locations. Indeed SDF-1 is chemoattractant for human CD34+ progenitor cells.

35 Aiuti, et al. (1997) <u>J. Exp. Med.</u> 185:111-120. The observation that TECK is chemoattractant for thymocytes but not for mature peripheral T cells suggests that TECK could attract T-cell progenitors to the thymus. Such populations are very difficult to isolate in sufficient

numbers to conduct in vitro chemotaxis experiments, but we are currently designing new strategies to address this important question. In addition, we have not found significant chemotactic activity of TECK on bone marrow cells. SDF-1 was shown to be much less potent on CD34+ progenitors from the peripheral blood than those from the bone marrow. Aiuti, et al. (1997). It is possible that the sensitivity of progenitor cells to TECK would increase as these cells leave the bone marrow to colonize lymphoid organs. Importantly, intra-thymic maturation is 10 also characterized by a directional migration from the subcapsular region which contains the earliest progenitors to the cortex and finally to the medulla where thymocytes finish their maturation (Boyd, et al. (1993)). It is possible that the secretion of TECK by medullary dendritic cells may play a role in this directional migration. Yet another possibility is that TECK may play a role in the organization and development of the thymic stroma.

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We also showed that TECK is chemotactic for activated macrophages and dendritic cells. These two cell types also play important roles in T-cell development. Through a complex screening process involving positive and negative selection events most of the antigenic specificities randomly generated in the thymus will be eliminated by programmed cell death (Janeway (1994) Immunity 1:3-6). The efficient scavenging of dead thymocytes is probably mediated, at least partially, by thymic macrophages and thus TECK could play an important role through its action on 30 activated macrophages. Further along, T-cells with a high affinity for self-antigens and thus potentially harmful are eliminated through negative selection (Janeway (1994)). It is believed that thymic dendritic cells are primarily responsible for the negative 35 selection of thymocytes, therefore playing a major role in the establishment of tolerance. Inaba, et al. (1991) J. Exp. Med. 173:549-559. An efficient mechanism of central tolerance should eliminate T cells potentially

reactive against auto-antigens which are not expressed in the thymus, such as organ specific auto-antigens. Several known chemokines induce the migration of dendritic cells and could therefore contribute to their recruitment during peripheral immune responses. Sozzani, et al. (1995) J. Immunol. 155:3292-3295; and Xu, et al. (1996) <u>J. Leukoc. Biol.</u> 60:365-371. Similarly, dendritic cells presenting organ-specific or other antigens could be recruited to the thymus or the small intestine and induce negative selection of T cells specific for these 10 antigens. It is possible that thymus- and small intestine-specific chemokines active on dendritic cells such as TECK could play an important role in the establishment of tolerance. Thus, TECK could potentially interact at several important steps of T-cell 15 development. Future experiments will aim to define the precise role of TECK in T-cell development and other physiological processes through the use of genetically modified mice.

20 I. TECK is specifically expressed by thymic dendritic cells

Dendritic cells represent a heterogeneous cell population derived from bone marrow progenitors. They are present in non-lymphoid organs as immature dendritic cells (such as Langerhans cells in the skin) where they 25 display a high ability for antigen capture. Cella, et al. (1997) Curr. Opin. Immunol. 9:10-16. Subsequent to antigen challenge, they will migrate to secondary lymphoid organs and will acquire a high capacity to present processed antigens to naive T-cells to initiate a 30 specific immune response (Cella, et al. (1997)). It has been shown that dendritic cells can derive from CD34+ progenitors cultured in the presence of GM-CSF and TNF- α (Caux, et al. (1992) Nature 360:258-261; and Caux, et al. (1996) J. Exp. Med. 184:695-706) or from monocytes in the presence of GM-CSF and IL-4 (Sallusto and Lanzavecchia (1994) J. Exp. Med. 179:1109-1118). Interestingly, there is also evidence for a lymphoid dendritic cell precursor in thymus and bone marrow which is able to derive both

lymphocytes and dendritic cells in the absence of GM-CSF. Ardavin, et al. (1993) Nature 362:761-763; Galy, et al. (1995) Immunity 3:459-473; Marquez, et al. (1995) J. Exp. Med. 181:475-483; and Wu, et al. (1996). These lymphoidderived dendritic cells may have different functional properties such as a negative regulation of T-cell responses since they express FasL in the mouse. Suss and Shortman (1996) J. Exp. Med. 183:1789-1796. We found that TECK was expressed at high levels in mouse thymic dendritic cells but was absent in cDNA libraries from 10 mouse splenic dendritic cells or from human dendritic cells generated in vitro from CD34+ precursors or monocytes. Interestingly, mTECK mRNA was present at a low level in a population of early thymocyte progenitors still able to derive dendritic cells (Wu, et al. (1996). Thus, it would be tempting to suggest that TECK could be a specific marker of lymphoid-derived dendritic cells. However, we observed that TECK was absent from splenic dendritic cells that likely contain lymphoid-derived dendritic cells. The expression of TECK mRNA appeared in 20 the spleen of mice injected with LPS would suggest that peripheral dendritic cells may express TECK upon activation, but we found that TECK was not expressed in cDNA libraries of bone-marrow derived dendritic cells activated with LPS, PMA and ionomycin or IL-1 α and TNF- α . 25 It is possible that the normal expression of TECK is specific for lymphoid-derived dendritic cells or, alternatively, that it is upregulated by very specific stimuli present in the thymic and intestinal microenvironment under physiological conditions. Consistent 30 with the latter hypothesis is our observation of specific staining of thymic endothelial cells with anti-TECK antibody since we have not been able to find TECK expression in human HUVEC endothelial cells by northern 35 blot analysis, without activation or following a 16 houractivation with various combinations of IL-1, TNF- α , IL-4, IL-7 and oncostatin while some of these stimuli induce the expression of other CC chemokines in endothelial cells. Rollins and Pober (1991) Am. J. Pathol. 138:13151319; Marfaing-Koka, et al. (1995) <u>J. Immunol.</u> 154:1870-1878; Garcia-Zepeda, et al. (1996) <u>J. Immunol.</u> 157:5613-5626; and Garcia-Zepeda, et al. (1996) <u>Nat. Med.</u> 4:449-456. Taken together, our data strongly suggest that TECK is a novel chemokine specifically expressed by activated lymphoid-derived dendritic cells.

Through their function of antigen presentation, dendritic cells play major roles in the establishment of tolerance and in the initiation of an antigen-specific immune response. The use of purified dendritic cells has been recently proposed in different therapeutic protocols (Cella, et al. (1997)). The discovery of factors with a regulated expression in dendritic cells such as the novel CC chemokine TECK will certainly improve our knowledge of the biology of dendritic cells and lead to the design of relevant in vivo applications.

J. Mice and in vivo experimental procedures

Four to eight week-old and time-pregnant BALB/c mice were purchased from Simonsen Laboratories (Gilroy, CA).

RAG-1-deficient mice (Mombaerts, et al. (1992)) were purchased from The Jackson Laboratory (Bar Harbor, ME).

To analyze TECK expression after in vivo activation, various organs were recovered from pools of 2 mice 3

hours after intravenous LPS injection (50 μ g LPS in 200

 μ 1 PBS or 200 μ 1 PBS for controls).

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K. Cell purification, culture and stimulation.

THP-1 cells (TIB-202 from the American Type Culture Collection, Rockville, MD) were cultured in complete medium which consisted in RPMI 1640 medium (JRH BioSciences, Lenexa, KS) supplemented with 10% FCS, 200 mM L-glutamin, 5 x 10⁻⁵ M mercaptoethanol, MEM aminoacids and vitamins, sodium bicarbonate, penicillin, streptomycin (all from Sigma, ST. Louis, MO), and gentamycin (Boehringer, Indianapolis, IN). To obtain activated mouse macrophages, 10 ml of cold PBS were injected into the peritoneum and the collected cells allowed to adhere to plastic for 24 h in complete medium. The adherent fraction, mostly macrophages, was then

collected. To obtain splenic dendritic cells, a

splenocyte cell suspension was prepared in RPMI 1640 Dutch modified medium (Life Technologies, Paisley, Scotland) as described previously in, e.g., Macatonia, et al. (1987) J. Exp. Med. 166:1654-1667. Splenocytes were incubated at 37°C for 16 h and the cell suspension was collected and laid over Metrizamide (Nycomed Pharma, Oslo, Norway). After centrifugation for 10 min. at 1700 x g, the low interface was collected and stained with anti-Mac-1 (Pharmingen, San Diego, CA) and the anti-CD11c N-418 antibodies (Macatonia, et al. (1993) J. Immunol. 10 150:3755-3765). Splenic dendritic cells were sorted by flow cytometry on a FACStar plus cell sorter (Becton Dickinson, Mountain View, CA) to a purity greater than 98% upon reanalysis in all the experiments included in this report. To obtain thymic dendritic cells, thymuses 15 were cut in small fragments and resuspended in 10 ml of RPMI-1640 +10% FCS containing 1 mg/ml collagenase and 0.02 mg/ml DNase I (both from SIGMA) and digested with continuous agitation at room temperature for 30 min. 20 (Shortman, et al. (1995) Adv. Exp. Med. Biol. 378:21-29). One ml of 0.1M EDTA pH 7.2 was added for an additional 5 min. Cells were then washed in complete medium, resuspended in complete medium and overlaid onto Metrizamide. The thymic dendritic cell-enriched preparation was then stained with anti-IAd and N-418 antibodies and the dendritic cells sorted by flow

L. Molecular cloning of mouse and human TECK

The cDNA encoding mouse TECK was obtained by random sequencing of a RAG-1 KO mouse thymic directional cDNA library. Briefly, mRNA was extracted using RNAzolTM B (Tel-Test, Friendswood, TX) and then oligotex-dT mRNA kit (Quiagen, Chatsworth, CA) following the manufacturer's instruction. A directional cDNA library was prepared using the SuperscriptTM Plasmid System (Gibco-BRL, Grand Island, NY) and cloned into the pME18s plasmid vector. Sequencing was done using the TaQ DyeDeoxy Terminator Cycle Sequencing kit (Applied Biosystems, Foster City, CA). To determine whether TECK was present in other

cytometry

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mammals including human, a Southern blot containing EcoRI digested genomic DNA from different species (Bios Laboratories, New Haven, CT) was hybridized with the full-length mouse TECK cDNA.

The cDNA encoding human TECK was found by screening of a small intestine cDNA library using the full-length mouse TECK cDNA as a probe following standard procedures.

M. Northern blot analysis of RNA and Southern blot analysis of cDNA libraries

10 All RNA's were isolated from tissues or cells using RNAzol $^{ extsf{TM}}$ B (Tel-Test) and analyzed after electrophoresis in a 1% formaldehyde-agarose gel (10 μ g/lane). RNA's were then blotted onto a Hybond-N+ nylon membrane (Amersham, Arlington Heights, IL). Some northern blots of mRNA were bought from Clontech (Palo Alto, CA). 15 analyze the expression of TECK in cDNA libraries (obtained from T. MacClanahan, DNAX), 10 μg of DNA were digested with the appropriate restriction enzymes to release their inserts and analyze by Southern blotting onto nylon membranes. Northern blots and blots of cDNA 20 libraries were hybridized for 16 hours at 65°C with a 32 P-labeled probe consisting in the full-length cDNA encoding for mouse or human TECK and then washed and exposed, according to standard protocols.

25 N. Inter specific mouse backcross mapping

Inter specific backcross progeny were generated by mating (C57B1/6J x M. spretus) F1 females and C57B1/6J males as described, e.g., in Copeland and Jenkins (1991) Trends Genet. 7:113-118. A total of 205 N2 mice was used to map the Teck locus. DNA isolation, restriction enzyme 30 digestion, agarose gel electrophoresis, Southern blot transfer and hybridization with the full-length mTECK cDNA probe were performed as described, e.g., in Jenkins, et al. (1982). Fragments of 7.5, 6.9, and 2.5 kb were 35 detected in HincII digested C57Bl/6J DNA and fragments of 8.8 and 5.4 kb were detected in HincII digested M. spretus DNA. The presence or absence of the 8.8 and 5.4 kb HincII M. spretus-specific fragments, which cosegregated, was followed in backcross mice. A

description of the probes and RFLPs for two of the loci linked to <u>Teck</u> including <u>Insr</u> has been reported previously, e.g., in Ceci, et al. (1990) <u>Genomics</u> 6:72-79. Recombination distances were calculated as described (Green (1981) "Linkage, recombination and mapping" pp. 77-113 in <u>Genetics and Probability in Animal Breeding</u> <u>Experiments</u>, Oxford University Press, New York) using the computer program SPRETUS MADNESS.

- O. Measurement of TECK mRNA expression by RT-PCR RNA's from sorted thymic dendritic cells or fetal 10 thymuses were prepared with the RNeasy total RNA kit (Quiagen, Chatsworth, CA), following the manufacturer's instructions. First strand cDNAs were generated by reverse transcription with a random hexamer in a 10 μ 1 reaction and 1 μ l of this reaction was used as a template 15 for PCR. TECK expression was compared to the expression of hypoxanthine-guanine phosphoribosyl transferase (HPRT). Primer sequences were as follows: TECK: 5' primer, 5'CCTTCAGGTATCTGGAGAGGAGATC3' (nucleotides 58-72 of SEQ ID NO: 1) and 3' primer, 20 5'CACGCTTGTACTGTTGGGGTTC3' (complement of nucleotides 447-468 of SEQ ID NO: 1), HPRT: 5' primer, 5'GTAATGATCAGTCAACGGGGGAC3' (SEQ ID NO: 17) and 3' primer, 5'CCAGCAAGCTTGCAACCTTAACCA3' (SEQ ID NO: 18). Samples were submitted to 25 cycles of amplification, 25 each composed of 94°C for 1 min., 57°C for 30 s and 72° C for 2 min. PCR products were then separated by
- Biotin-14-CTP labeled sense and antisense riboprobes were generated using a non radioactive RNA labeling system (Gibco, Gaithersburg, MD) and the plasmid PCRII (InVitrogen, Carlsbad, CA) containing a 400 base pair TECK cDNA fragment inserted by PCR and TA cloning (InVitrogen). Paraffin-embedded tissues were cut in 3-5 µm sections, mounted on slides, baked at 60°C for one hour, deparaffinized in xylene (Fisher Scientific, Pittsburgh, PA) and immersed in 100% ethanol. Sections

electrophoresis in 2% agarose gels and stained with

ethidium bromide.

were then incubated for 10 min at 37°C in proteinase K solution (40 mg/ml) (Gibco) in PBS and rinsed for 2 min in PBS at room temperature before being refixed in 10% formalin (Fisher Scientific, Pittsburgh, PA) in PBS for 1 min. Next, the sections were dehydrated through graded solutions of ethanol and air dried. Hybridization was carried out using the Gibco in situ hybridization and detection system kit. Vanadyl ribonucleoside complex (Gibco) was added to the hybridization solution (39 mM 10 final). A 0.1 μ g/ml concentration of each probe was used during an 18 h hybridization at 42° C. Posthybridization washes used room temperature 0.2% SSC. Following detection and substrate visualization, the slides were counterstained with 1% nuclear red stain 15 (Sigma, St. Louis, MO).

Q. Immunohistochemistry

A polyclonal antibody specific of a synthetic decapeptide identical to the C-terminus part of murine TECK (Figure 1) was prepared in rabbits by Research Genetics (Huntsville, AL). Normal rabbit serum from a 20 pool of 50 different animals (Research Genetics) was used as a negative control. To study TECK protein expression in the mouse thymus, 6 µm thick cryostat sections were thaw mounted on organosilicone subbed slides (American 25 Histology Reagent Co., Stockton, CA.) and fixed in 3% formalin (Fisher Scientific, Springfield, NJ) in Hank's Balanced Salt Solution with 0.01M HEPES (HBSS-HEPES), pH 7.4, for 15 min at room temperature. The sections were sequentially blocked for endogenous biotin binding using the Vector blocking kit (Vector Laboratories, Burlingame, CA) and for endogenous peroxidase activity with a 1% hydrogen peroxide, 0.2M sodium azide solution, in HBSS-HEPES with 0.1% saponin (staining buffer). Non-specific antibody binding sites were then blocked with 10% normal goat serum (Sigma) in staining buffer. Sections prepared 35 as above were first incubated for 18 h at 25°C with 1/500 dilution of polyclonal antibody or control rabbit serum in staining buffer. In the second step, the sections were incubated for 1 h at room temperature with biotin

labeled goat anti-rabbit IgG (2 μg/ml) (Vector Laboratories) in staining buffer and then for 30 min at room temperature with the Vectastain Elite ABC Kit (Vector Laboratories) in staining buffer. The sections were then rinsed in HBSS-HEPES without saponin.

Immunoenzyme tissue staining was revealed with 3, 3' - diaminobenzidine tetrahydrochloride (DAB) substrate (0.5 mg/ml) (Sigma) in 0.05M Tris, pH 7.4, containing 0.0075% hydrogen peroxide. The substrate reaction was stopped by rinsing the sections in distilled water. The sections were then counterstained with Harris' hematoxylin (Shandon Lipshaw, Pittsburg, PA).

The expression of TECK mRNA in murine adult thymus was analyzed by in situ hybridization and revealed a discrete positive non-lymphoid population within the thymus medulla. The expression of TECK protein was analyzed by using a polyclonal anti-serum made in a rabbit immunised with a peptide that consisted in the last 12 amino-acid of the murine TECK protein sequence. This polyclonal antibody reacts with the murine TECK recombinant protein prepared at DNAX both in ELISA and western blot. The application of this anti-serum on mouse adult thymic sections confirmed the distribution pattern obtained by in situ hybridization: the cells producing TECK are medullary stromal cells. The precise cell type producing TECK within the mouse thymus was identified, using the same anti-serum on sorted thymic subsets, as being the thymic dendritic cells. R. Production of recombinant mouse TECK in Escherichia coli and other chemokines

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Mouse recombinant TECK was produced in E. coli as an N-terminal FLAG (DYKDDDDKL; SEQ ID NO: 19) fusion protein. Briefly, the fusion construct containing FLAG followed by the mTECK sequence minus the leader peptide (see SEQ ID NO: 2) was obtained by PCR amplification of the TECK cDNA in order to flank the coding sequence with HindIII and EcoRI sites and subsequent ligation in the pFLAG.1 vector which contains the FLAG sequence and an OmpA signal sequence. Electro-competent UT 4400 E. coli

were transformed with the pFLAG.1-mTECK plasmid. cells were grown in 2 x LB plus 50 µg/ml Ampicillin, induced at an OD. of 2.3 with 400 μM IPTG and harvested. The cell pellet was resuspended in cold lysis buffer (20 mM Tris pH 8, 2 mM EDTA, 20% sucrose, 0.1 mg/ml lysozyme. 100 μ l Benzonase), homogenized and allowed to sit for 30 Then the same amount of a 1:4 dilution of cold lysis buffer without lysozyme was added for 10 more min. The solution was spun and the supernatant was filtered 10 through a 0.2 μm membrane and then diluted 1:1 in 50 mMTris pH 7.5. The diluted osmotic extract was submitted to chromatography on a Q-sepharose column equilibrated with 50 mM Tris pH 7.5 and eluted with a linear salt gradient. The fractions containing the recombinant 15 protein were pooled. The fractions were then loaded onto a S-sepharose column equilibrated with 20 mM acetate pH The column was eluted with a linear salt gradient and then with a 1.5M NaCl wash that contained the protein. Finally, the eluate was loaded onto a reverse phase column. The column was eluted with a linear 20 gradient of 20% to 80% acetonitrile + 0.1% TFA. concentration of the mTECK protein was estimated by Comassie blue staining and densitometric scanning of a 10% Nu-PAGE gel with lysozyme as a standard. The purity was estimated at 100% by sequencing of the N-terminus of 25 the recombinant protein. Recombinant murine MIP-1 α (R&D Systems, Minneapolis, MN) and lymphotactin (Hedrick, et al. (1997) J. Immunol. 158:1533-1540) were used as controls.

30 S. Assay for chemotaxis

The in vitro migration of cells isolated as described above in response to TECK or other factors was assessed in a modified Boyden micro chamber (Neuroprobe, Cabin John, MD) as described previously (Kelner, et al.

35 (1994)). Briefly, factor dilutions in DMEM medium (Gibco) were loaded in the lower compartment in duplicate and 10^5 cells in a 50 μl volume of DMEM were loaded in the upper compartment. The two compartments were separated by a 5- μm or 8- μm pore size polycarbonate

filter (Nucleopore, Pleasanton, CA). After incubation at 37°C for 80 min (or 120 min for lymphocytes), the filters were fixed in methanol and stained with Fields A and B. Cell migrated on the other side of the membrane were counted per five high-power fields (100 x) under microscope. The chemotactic index was calculated from the number of cells counted with the test sample divided by the number of cells counted with medium alone.

Northern blot analysis was performed of RNA from different organs hybridized with the mTECK cDNA probe with or without in vivo LPS stimulation. Hybridizing bands corresponded to the predicted =1040 bp size for mTECK mRNA. Significant induction occurred in spleen (with virtually no background), and in thymus and small intestine (both with higher background); no signal was detected in either condition for heart, lung, kidney, or liver.

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mTECK mRNA expression was analysed in the mouse fetal thymus. RNA's from fetal thymic lobes were extracted at day 14, 15, 16 and 17 of gestation. Positive RT-PCR signals were detected in each of day 14, 125, 16, and 17 samples.

mTECK mRNA expression in thymic dendritic cells was evaluated. A population enriched in thymic dendritic cells was prepared from 15 pooled adult thymuses. >99% pure dendritic cells were then sorted by flow cytometry based on their MHC Class II+ N-418+ phenotype. mTECK mRNA was then analyzed by RT-PCR and a MHC class II+ N-418- population sorted in the same experiment was used as a negative control. The N418+ sample gave a positive signal, while the N418- sample did not.

Expression analysis was performed with hTECK mRNA in different Human Tissues and Cell Types. Southern blots of human cDNA libraries digested with the appropriate restriction enzymes were hybridized with the hTECK cDNA probe. A major band hybridizing corresponding to the predicted length of hTECK mRNA (=1040 bp) was observed with sometimes some other bands that may represent incomplete cDNAs. Positive signals were detected in

tonsil, fetal spleen, and fetal small intestine. No signal was detected in activated (with PMA and ionomycin for 12 h) NK cells, activated (anti-CD40 antibody and IL-4 for 6 and 12 h) splenocytes, $\gamma\delta$ T cells, activated

- (with anti-CD3 and PMA for 6, 12, and 24 h) PBMC, fetal testis, C+ (elutriated monocytes cultured with IFN-γ and IL-10) monocytes, C- monocytes, 70% pure dendritic cells (CD1α+ dendritic cell population obtained by expansion of CD34+ bone marrow cells with GM-CSF and TNF-α and
- resting), and DC3 (similar dendritic cell population stimulated with PMA and ionomycin for 1 and 6 h), DC5 (dendritic cells obtained by culturing peripheral blood monocytes in the presence of IL-4 and GM-CSF), U937 (premonocytic cell line), and CD1α cell lines. Ras KO
- mouse cDNA again confirmed that the mouse and human genes crosshybridize.

Four independent lines of transgenic mice expressing TECK in the brain have been made. All animals had neurologic disorders. In addition, several of them suffered severe infections. The consequences of TECK could be a direct one on brain cells which nature remains to be identify. Alternatively, since TECK has been shown in vitro to have effects on macrophages and dendritic cells which are critical effectors of immune responses, the overproduction of TECK could lead to distant effects on these cells at sites of infection. These results

suggest that the blockade of TECK production in vivo may help to resolve particular pathological processes, in

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VII. Screening for receptor/ligand

particular infections.

Labeled reagent is useful for screening of an expression library made from a cell line which expresses a chemokine or receptor, as appropriate. Standard staining techniques are used to detect or sort intracellular or surface expressed ligand, or surface expressing transformed cells are screened by panning. Screening of intracellular expression is performed by

various staining or immunofluorescence procedures. See also, e.g., McMahan, et al. (1991) EMBO J. 10:2821-2832.

For example, on day 0, precoat 2-chamber permanox slides with 1 ml per chamber of fibronectin, 10 ng/ml in PBS, for 30 min at room temperature. Rinse once with PBS. Then plate COS cells at 2-3 x 10⁵ cells per chamber in 1.5 ml of growth media. Incubate overnight at 37° C.

On day 1 for each sample, prepare 0.5 ml of a solution of 66 µg/ml DEAE-dextran, 66 µM chloroquine, and 4 µg DNA in serum free DME. For each set, a positive control is prepared, e.g., of huIL-10-FLAG cDNA at 1 and 1/200 dilution, and a negative mock. Rinse cells with serum free DME. Add the DNA solution and incubate 5 hr at 37° C. Remove the medium and add 0.5 ml 10% DMSO in DME for 2.5 min. Remove and wash once with DME. Add 1.5 ml growth medium and incubate overnight.

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On day 2, change the medium. On days 3 or 4, the cells are fixed and stained. Rinse the cells twice with Hank's Buffered Saline Solution (HBSS) and fix in 4% 20 paraformaldehyde (PFA)/glucose for 5 min. Wash 3X with The slides may be stored at -80° C after all liquid is removed. For each chamber, 0.5 ml incubations are performed as follows. Add HBSS/saponin(0.1%) with 32 μ l/ml of 1M NaN3 for 20 min. Cells are then washed with HBSS/saponin 1X. Add antibody complex to cells and 25 incubate for 30 min. Wash cells twice with HBSS/saponin. Add second antibody, e.g., Vector anti-mouse antibody, at 1/200 dilution, and incubate for 30 min. Prepare ELISA solution, e.g., Vector Elite ABC horseradish peroxidase solution, and preincubate for 30 min. Use, e.g., 1 drop 30 of solution A (avidin) and 1 drop solution B (biotin) per 2.5 ml HBSS/saponin. Wash cells twice with HBSS/saponin. Add ABC HRP solution and incubate for 30 min. Wash cells twice with HBSS, second wash for 2 min, which closes cells. Then add Vector diaminobenzoic acid (DAB) for 5 35 to 10 min. Use 2 drops of buffer plus 4 drops DAB plus 2 drops of H_2O_2 per 5 ml of glass distilled water. Carefully remove chamber and rinse slide in water. Air

dry for a few minutes, then add 1 drop of Crystal Mount and a cover slip. Bake for 5 min at 85-90° C.

Alternatively, the binding compositions are used to affinity purify or sort out cells expressing the ligand or receptor. See, e.g., Sambrook, et al. or Ausubel et al.

All references cited herein are incorporated herein by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

10

SEQUENCE LISTING

-	(1) GENERAL INFORMATION:	
5	(i) APPLICANT: (A) NAME: Schering Corp.	
10	(B) STREET: 2000 Galloping Hill Road (C) CITY: Kenilworth (D) STATE: New Jersey (E) COUNTRY: USA (F) ZIP: 07033-0530 (G) TELEPHONE: 908-298-2906 (H) TELEFAX: 908-298-5388	
15		
	(ii) TITLE OF INVENTION: MAMMALIAN CHEMOKINE REAGENTS	•
20	(iii) NUMBER OF SEQUENCES: 19	
25	(iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: Apple Macintosh (C) OPERATING SYSTEM: Macintosh 7.1 (D) SOFTWARE: Microsoft Word 5.1a	
30	(v) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: (C) CLASSIFICATION:	
35	(vi) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 08/675,814 (B) FILING DATE: 05-JUL-1996	
40	(2) INFORMATION FOR SEQ ID NO:1: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1034 base pairs	
	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
45	(ii) MOLECULE TYPE: cDNA	-
50	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 94525	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
55		60
	TCAGGTATCT GGAGAGGAGA TCTAACCTTC ACT ATG AAA CTG TGG CTT TTT GCC Met Lys Leu Trp Leu Phe Ala 1 5	114
60	TGC CTG GTT GCC TGT TTT GTT GGG GCC TGG ATG CCG GTT GTC CAT GCC Cys Leu Val Ala Cys Phe Val Gly Ala Trp Met Pro Val Val His Ala 10 15 20	162

	CAA	GGT	GCC	TTT	GAA	GAC	TGC	TGC	CTG	GGT	TAC	CAG	CAC	AGG	ATC	AAA	210
	Gln	Gly 25	Ala	Phe	Glu	Asp	Cys 30	Суз	Leu	Gly	Tyr	Gln 35	His	Arg	Ile	Lys	
5	TGG Trp 40	AAT Asn	GTT Val	CTC	CGG Arg	CAT His 45	GCT Ala	AGG Arg	AAT Asn	TAT Tyr	CAC His 50	Gln	CAG Gln	GAA Glu	GTG Val	AGT Ser 55	258
10	GGA Gly	AGC Ser	TGC Cys	AAC Asn	CTA Leu 60	CGT Arg	GCT Ala	GTG Val	AGA Arg	TTC Phe 65	TAC Tyr	TTC Phe	CGC Arg	CAG Gln	AAA Lys 70	GTA Val	306
15	GTG Val	TGT Cys	GGG Gly	AAT Asn 75	CCA Pro	GAG Glu	GAC Asp	ATG Met	AAT Asn 80	GTG Val	AAG Lys	AGG Arg	GCG Ala	ATA Ile 85	AGA Arg	ATC Ile	354
20	TTG Leu	ACA Thr	GCT Ala 90	AGG Arg	AAA Lys	AGG Arg	CTA Leu	GTC Val 95	CAC His	TGG Trp	AAG Lys	AGC Ser	GCC Ala 100	TCA Ser	GAC Asp	TCT Ser	402
20	CAG Gln	ACT Thr 105	GAA Glu	AGG Arg	AAG Lys	AAG Lys	TCA Ser 110	AAC Asn	CAT His	ATG Met	AAG Lys	TCC Ser 115	AAG Lys	GTG Val	GAG Glu	AAC Asn	450
25	CCC Pro 120	AAC Asn	AGT Ser	ACA Thr	AGC Ser	GTG Val 125	AGG Arg	AGT Ser	GCC Ala	ACC Thr	CTA Leu 130	GGT Gly	CAT His	CCC Pro	AGG Arg	ATG Met 135	498
30	GTG . Val	ATG Met	ATG Met	CCC Pro	AGA Arg 140	AAG Lys	ACC Thr	AAC Asn	AAT Asn	TAAG	TTA.	TT A	CTCA	GAG1	'A		545
	AGCA	CCAG	CT G	GAGG	ATGG	G CG	GAGT	CTGC	TGA	AGTG	CTG	тстт	CTAG	GC A	TGCC	AGTGC	605
35																CAGCA	665
																GGCTA	725
40																GAGTA	785
																ATTAA	845
45																GTTAG	905
40																GCCTG	965
	AAAA			TGGT	CAAA	G AA	AATT	AAAA	TAA	AAAC	TTA	AAAA	.GCTA	TT A	AAAA	GTAAA	1025
50	winn	MALA	ur.														1034

(2) INFORMATION FOR SEQ ID NO:2:

55

60

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 144 amino acids

- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Lys Leu Trp Leu Phe Ala Cys Leu Val Ala Cys Phe Val Gly Ala 1 5 15

	Trp	Met	Pro	Val 20	Val	His	Ala	Gln	Gly 25	Ala	Phe	Glu	Asp	Cys 30	Суѕ	Leu	1		
5	Gly	Tyr	Gln 35	His	Arg	Ile	Lys	Trp 40	Asn	Val	Leu	Arg	His 45	Ala	Arg	Asn	1		
	Tyr	His 50	Gln	Gln	Glu	Val	Ser 55	Gly	Ser	Cys	Asn	Leu 60	Arg	Ala	Val	Arg	ı		
LO	Phe 65	Tyr	Phe	Arg	Gln	Lys 70	Val	Val	Cys	Gly	Asn 75	Pro	Glu	Àsp	Met	Asn 80))		
15	Val	Lys	Arg	Ala	Ile 85	Arg	Ile	Leu	Thr	Ala 90	Arg	Lys	Arg	Leu	Val 95	His	3		
	Trp	Lys	Ser	Ala 100	Ser	Asp	Ser	Gln	Thr 105	Glu	Arg	Lys	Lys	Ser 110	Asn	His	3		
20	Met	Lys	Ser 115		Val	Glu	Åsn	Pro 120	Asn	Ser	Thr	Ser	Val 125	Arg	Ser	Ala	3		
25	Thr	Leu 130		His	Pro	Arg	Met 135	Val	Met	Met	Pro	Arg 140	Lys	Thr	Asn	Ası	n.		
	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:3	:										
	(2)) SE																
30		(1	(A) L B) T C) S	ENGT YPE: TRAN	H: 1 nuc	012 leic ESS:	base aci sir	pai .d	rs.									
35		(ii	L) MC	LECU	ILE T	YPE:	cDN	IA				•							
40		(i)		(A) 1	JAME/	KEY:		5 75(56										
		(iz	k) FI	(A) I	NAME	KEY	: mai	c_pe	ptide 66	e					,				
45	•								,										
			i) S																
	TC	GACC	CACG	CGT	CCGC	TTG (GCCT	ACAG	cc c	GGCG	GGCA'	T CA	GCTC	CCTT	GAC	CCAG	TGG	6	50
50	AT.	ATCG	GTGG	ccc	CGTT	ATT	CGTC	CAGG	TG C	CCAG	GGAG	G AG	SACC	CGCC	TGC	AGC		11	. 6
55	AT Me -2	t As	C CT n Le	G TG u Tr -2	p Le	C CT u Le	G GC	C TG a Cy	C CT s Le -1	u Va	G GC 1 Al	C GG a Gl	C TT y Ph	C CT e Le -1	u Gr	A GO y Al	CC la	16	54
	TG Tr	G GC	a Pr	C GC	T GI .a Va	C CA	C AC	C CA	A GC n Gl	T GT y Va	C TT	T GA Le Gl	G GA u As 5	C TC	C TG	C C	rG eu	2:	12
60	A)	C TA	AC CA /r Hi	C TA	AC CC	o I	TT GO	G TO Ly Ti	G GC	T GI la Va	ll Le	C CG	G CG	G GC g Al	C TG	P I	CT hr 25	2	6(

	TAC CGG ATC CAG GAG GTG AGC GGG AGC TGC AAT CTG CCT GCT GCG ATA Tyr Arg Ile Gln Glu Val Ser Gly Ser Cys Asn Leu Pro Ala Ala Ile 30 35 40	308
5	TTC TAC CTC CCC AAG AGA CAC AGG AAG GTG TGT GGG AAC CCC AAA AGC Phe Tyr Leu Pro Lys Arg His Arg Lys Val Cys Gly Asn Pro Lys Ser 45 50 55	356
10 ·	AGG GAG GTG CAG AGA GCC ATG AAG CTC CTG GAT GCT CGA AAT AAG GTT Arg Glu Val Gln Arg Ala Met Lys Leu Leu Asp Ala Arg Asn Lys Val 60 65 70	404
15	TTT GCA AAG CTC CAC CAC AAC ATG CAG ACC TTC CAA GCA GGC CCT CAT Phe Ala Lys Leu His His Asn Met Gln Thr Phe Gln Ala Gly Pro His 75 80 85	452
20	GCT GTA AAG AAG TTG AGT TCT GGA AAC TCC AAG TTA TCA TCA TCC AAG Ala Val Lys Lys Leu Ser Ser Gly Asn Ser Lys Leu Ser Ser Ser Lys 90 95 100 105	500
	TTT AGC AAT CCC ATC AGC AGC AGC AAG AGG AAT GTC TCC CTC CTG ATA Phe Ser Asn Pro Ile Ser Ser Ser Lys Arg Asn Val Ser Leu Leu Ile 110 115 120	548
25	TCA GCT AAT TCA GGA CTG TGAGCCGGCT CATTTCTGGG CTCCATCGGC Ser Ala Asn Ser Gly Leu 125	596
30	ACAGGAGGG CCGGATCTTT CTCCGATAAA ACCGTCGCCC TACAGACCCA GCTGTCCCCA	656
30	CGCCTCTGTC TTTTGGGTCA AGTCTTAATC CCTGCACCTG AGTTGGTCCT CCCTCTGCAC	716
	CCCCACCACC TCCTGCCCGT CTGGCAACTG GAAAGAAGGA GTTGGCCTGA TTTTAACCTT	776
35	TTGCCGCTCC GGGGAACAGC ACAATCCTGG GCAGCCAGTG GCTCTTGTAG AGAAAACTTA	836
	GGATACCTCT CTCACTTTCT GTTTCTTGCC GTCCACCCCG GGCCATGCCA GTGTGTCCTC	896
40	TGGGTCCCCT CCAAAAATCT GGTCATTCAA GGATCCCCTC CCAAGGCTAT GCTTTTCTAT	956
40	AACTITTAAA TAAACCTTGG GGGGTGAATG GAATAAAAAA AAAAAAAAA AAAAAA	1012
45	(2) INFORMATION FOR SEQ ID NO:4: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 150 amino acids	
50	(B) TYPE: amino acid (D) TOPOLOGY: linear	
50	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
55	Met Asn Leu Trp Leu Leu Ala Cys Leu Val Ala Gly Phe Leu Gly Ala -23 -15 -10	
60	Trp Ala Pro Ala Val His Thr Gln Gly Val Phe Glu Asp Cys Cys Leu -5 1 5	
	Ala Tyr His Tyr Pro Ile Gly Trp Ala Val Leu Arg Arg Ala Trp Thr 10 20 25	

	Tyr	Arg	Ile	Gln	Glu 30	Val	Ser	Gly	Ser	Cys 35	Asn	Leu	Pro	Ala	Ala 40	Ile	
5	Phe	Tyr	Leu	Pro 45	Lys	Arg	His	Arg	Lys 50	Val	Cys	Gly	Asn	Pro 55	Lys	Ser	
	Arg	Glu	Val 60	Gln	Arg	Ala	Met	Lys 65	Leu	Leu	Asp	Ala	Arg 70	Asn	Lys	Val	
10	Phe	Ala 75	Lys	Leu	His	His	Asn 80	Met	Gln	Thr	Phe	Gln 85	Ala	Gly	Pro	His	
15	Ala 90	Val	Lys	Lys	Leu	Ser 95	Ser	Gly	Asn	Ser	Lys 100	Leu	Ser	Ser	Ser	Lys 105	
	Phe	Ser	Asn	Pro	Ile 110	Ser	Ser	Ser	Lys	Arg 115	Asn	Val	Ser	Leu	Leu 120	Ile	
20	Ser	Ala	Asn	Ser 125	Gly	Leu											
	(2)						ID N										
25		(i)	I) I). I).	A) LI 3) Ti C) Si	engti Pe: Prani	nuc DEDNI	TERI 1 ba leic ESS: line	acio sing	pairs 1	3							
30		(ii)	MOI	LECUI	LE T	YPE:	cDNZ	\									
35		(ix)		A) N	AME/	KEY: ION:	CDS	288						•			
40			C	A) N	AME/I	ION:	mat_ 79.	. 288									
										ID N							
45	Met	TGC Cys -25	TGT Cys	ACC	AAG Lys	AGT Ser	TTG Leu -20	CTC Leu	CTG Leu	GCT Ala	GCT Ala	Leu -15	ATG Met	TCA Ser	Val	CTG Leu	48
50	CTA Leu -10	CTC Leu	CAC	CTC Leu	TGC Cys	GGC Gly -5	Glu	TCA Ser	GAA Glu	GCA Ala	GCA Ala 1	AGC Ser	AAC Asn	TTT Phe	GAC Asp 5	Суз	96
	TGT Cys	CTT Leu	GGA Gly	TAC Tyr 10	Thr	GAC Asp	CGT	ATT	CTT Leu 15	His	CCT Pro	AAA Lys	TTT Phe	ATT Ile .20	GTG Val	GGC	144
55	TTC Phe	ACA Thr	CGG Arg 25	Gln	CTG Leu	GCC Ala	AAT Asn	GAA Glu 30	Gly	TGT Cys	GAC Asp	ATC Ile	AAT Asn 35	Ala	ATC	ATC	192
60	TTT Phe	CAC His	Thr	AAG Lys	AAA Lys	AAG Lys	TTG Leu 45	Ser	GTG Val	TGC Cys	GCA Ala	AAT Asn 50	Pro	AAA Lys	CAG Gln	ACT Thr	240

	TIGG GTG AAA TAT ATT GTG CGT CTC CTC AGT AAA AAA GTC AAG AAC ATG Trp Val Lys Tyr Ile Val Arg Leu Leu Ser Lys Lys Val Lys Asn Met 55 60 65 70
5	TAAAAACTGT GGCTTTTCTG GAATGGAATT GGACATAGCC CAAGAACAGA AAGAACCTTG
	CTGGGGTTGG AGGTTTCACT TGCACATCAT GGAGGGTTTA GTGCTTATCT AATTTGTGCC
10	TCACTGGACT TGTCCAATTA ATGAAGTTGA TTCATATTGC ATCATAGTTT GCTTTGTTTA
	AGCATCACAT TAAAGTTAAA CTGTATTTTA TGTTATTTAT AGCTGTAGGT TTTCTGTGTT
	TAGCTATTTA ATACTAATTT TCCATAAGCT ATTTTGGTTT AGTGCAAAGT ATAAAATTAT
15 ,	ATTTGGGGGG GAATAAGATT ATATGGACTT TTTTGCAAGC AACAAGCTAT TTTTTAAAAA
`	AAACTATTTA ACATTCTTTT GTTTATATTG TTTTGTCTCC TAAATTGTTG TAATTGCATT
20	ATAAAATAAG AAAAATATTA ATAAGACAAA TATTGAAAAT AAAGAAACAA AAAGTTAAAA
	AAA AAAAAAAA AAAAAAAA AAAAAAAAA
25	(2) INFORMATION FOR SEQ ID NO:6:
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear
30	(ii) MOLECULE TYPE: protein
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
35	Met Cys Cys Thr Lys Ser Leu Leu Leu Ala Ala Leu Met Ser Val Leu -26 -25 -20 -15
40	Leu Leu His Leu Cys Gly Glu Ser Glu Ala Ala Ser Asn Phe Asp Cys -10 -5 1 5
40	Cys Leu Gly Tyr Thr Asp Arg Ile Leu His Pro Lys Phe Ile Val Gly 10 15 20
45	Phe Thr Arg Gln Leu Ala Asn Glu Gly Cys Asp Ile Asn Ala Ile Ile 25 30 35
	Phe His Thr Lys Lys Leu Ser Val Cys Ala Asn Pro Lys Gln Thr 40 45 50
50	Trp Val Lys Tyr Ile Val Arg Leu Leu Ser Lys Lys Val Lys Asn Met 60 65 70
55	(2) INFORMATION FOR SEQ ID NO:7:
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 699 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single
60	(D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

(A) NAME/KEY: CDS

	(B) LOCATION: 142435	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
	GGCACGAGCG GCACGAGCAT CACTCACACC TTGCATTTCA CCCCTGCATC CCAGTCGCCC	60
10	TGCAGCCTCA CACAGATCCT GCACACACCC AGACAGCTGG CGCTCACACA TTCACCGTTG	120
_	GCCTGCCTCT GTTCACCCTC C ATG GCC CTG CTA CTG GCC CTC AGC CTG CTG	171
	Met Ala Leu Leu Ala Leu Ser Leu Leu 1 5 10	
15	GTT CTC TGG ACT TCC CCA GCC CCA ACT CTG AGT GGC ACC AAT GAT GCT Val Leu Trp Thr Ser Pro Ala Pro Thr Leu Ser Gly Thr Asn Asp Ala 15 20 25	219
20	GAA GAC TGC TGC CTG TCT GTG ACC CAG AAA CCC ATC CCT GGG TAC ATC Glu Asp Cys Cys Leu Ser Val Thr Gln Lys Pro Ile Pro Gly Tyr Ile 30 35 40	267
25	GTG AGG AAC TTC CAC TAC CTT CTC ATC AAG GAT GGC TGC AGG GTG CCT Val Arg Asn Phe His Tyr Leu Leu Ile Lys Asp Gly Cys Arg Val Pro 45 50 55	315
30	GCT GTA GTG TTC ACC ACA CTG AGG GGC CGC CAG CTC TGT GCA CCC CCA Ala Val Val Phe Thr Thr Leu Arg Gly Arg Gln Leu Cys Ala Pro Pro 60 65 70	363
25	GAC CAG CCC TGG GTA GAA CGC ATC ATC CAG AGA CTG CAG AGG ACC TCA Asp Gln Pro Trp Val Glu Arg Ile Ile Gln Arg Leu Gln Arg Thr Ser 75 80 85 90	411
35	GCC AAG ATG AAG CGC CGC AGC AGT TAACCTATGA CCGTGCAGAG GGAGCCCGGA Ala Lys Met Lys Arg Arg Ser Ser 95	465
40	GTCCGAGTCA AGCATTGTGA ATTATTACCT AACCTGGGGA ACCGAGGACC AGAAGGAAGG	525
	ACCAGGCTTC CAGCTCCTCT GCACCAGACC TGACCAGCCA GGACAGGCCC TGGGGTGTGT	585
45	GTGAGTGTGA GTGTGAGCGA GAGGGTGAGT GTGGTCTAGA GTAAAGCTGC TCCACCCCCA GATTGCAATG CTACCAATAA AGCCGCCTGG TGTTTACAAC TAAAAAAAAA AAAA	64! 69!
50	(2) INFORMATION FOR SEQ ID NO:8: (i) SEQUENCE CHARACTERISTICS:	
55	(A) LENGTH: 98 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear	
J.J	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
60	Met Ala Leu Leu Ala Leu Ser Leu Leu Val Leu Trp Thr Ser Pro 1 5 10 - 15	
	Ala Pro Thr Leu Ser Gly Thr Asn Asp Ala Glu Asp Cys Cys Leu Ser 20 25 30	

	Val Thr Gln Lys Pro Ile Pro Gly Tyr Ile Val Arg Asn Phe His Tyr 35 40 45	
. 5	Leu Leu Ile Lys Asp Gly Cys Arg Val Pro Ala Val Val Phe Thr Thr 50 55 60	
10	Leu Arg Gly Arg Gln Leu Cys Ala Pro Pro Asp Gln Pro Trp Val Glu 65 70 75 80	
	Arg Ile Ile Gln Arg Leu Gln Arg Thr Ser Ala Lys Met Lys Arg Arg 85 90 95	
15	Ser Ser	
20	(2) INFORMATION FOR SEQ ID NO:9: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1119 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: CDNA	
30	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 11095 (D) OTHER INFORMATION: Nucleotide 579 may be A, C, G or T, and the codon may code for His or Gln.	
	·	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
35 40	ATG TTT TCG ACT CCA CTC AAC AND AND TOO TCG	48
40	ATG TTT TCG ACT CCA GTG AAG ATT ATT TTG TGT CAG TCA ATA CTT CAT Met Phe Ser Thr Pro Val Lys Ile Ile Leu Cys Gln Ser Ile Leu His 1 5 10 15	4 8 96
	ATG TTT TCG ACT CCA GTG AAG ATT ATT TTG TGT CAG TCA ATA CTT CAT Met Phe Ser Thr Pro Val Lys Ile Ile Leu Cys Gln Ser Ile Leu His 1	
40	ATG TTT TCG ACT CCA GTG AAG ATT ATT TTG TGT CAG TCA ATA CTT CAT Met Phe Ser Thr Pro Val Lys Ile Ile Leu Cys Gln Ser Ile Leu His 1	96
4 0	ATG TTT TCG ACT CCA GTG AAG ATT ATT TTG TGT CAG TCA ATA CTT CAT Met Phe Ser Thr Pro Val Lys Ile Ile Leu Cys Gln Ser Ile Leu His 1	96 44
4 0 4 5	ATG TTT TCG ACT CCA GTG AAG ATT ATT TTG TGT CAG TCA ATA CTT CAT Met Phe Ser Thr Pro Val Lys Ile Ile Leu Cys Gln Ser Ile Leu His 1	96 44

5	AAA Lys	GGC Gly	ATC Ile 115	TAT Tyr	GCC Ala	ATC Ile	AAC Asn	TTT Phe 120	AAC Asn	TGC Cys	GGG Gly	ATG Met	CTG Leu 125	CTC Leu	CTG Leu	ACT Thr	384
3	TGC Cys	ATT Ile 130	AGC Ser	ATG Met	GAC Asp	CGG Arg	TAC Tyr 135	ATC Ile	GCC Ala	ATT	Val	CAG Gln 140	GCG Ala	ACT Thr	AAG Lys	TCA Ser	432
10	TTC Phe 145	CGG Arg	CTC Leu	CGA Arg	TCC Ser	AGA Arg 150	ACA Thr	CTA Leu	CCG Pro	CGC	AGC Ser 155	AAA Lys	ATC Ile	ATC Ile	TGC Cys	CTT Leu 160	480
15	GTT Val	GTG Val	TGG Trp	GGG Gly	CTG Leu 165	TCA Ser	GTC Val	ATC Ile	ATC Ile	TCC Ser 170	AGC Ser	TCA Ser	ACT Thr	TTT Phe	GTC Val 175	TTC Phe	528
20	AAC Asn	CAA Gln	AAA Lys	TAC Tyr 180	AAC Asn	ACC Thr	CAA Gln	GGC Gly	AGC Ser 185	GAT Asp	GTC Val	TGT Cys	GAA Glu	CCC Pro 190	AAG Lys	TAC Tyr	576
25	CAN Xaa	ACT Thr	GTC Val 195	TCG Ser	GAG Glu	CCC Pro	ATC Ile	AGG Arg 200	TGG Trp	AAG Lys	CTG Leu	CTG Leu	ATG Met 205	TTG Leu	GGG	CTT Leu	624
	GAG Glu	CTA Leu 210	CTC Leu	TTT Phe	GGT Gly	TTC Phe	TTT Phe 215	ATC Ile	CCT Pro	TTG Leu	ATG Met	TTC Phe 220	ATG Met	ATA Ile	TTT Phe	TGT Cys	672
30	TAC Tyr 225	ACG Thr	TTC Phe	ATT Ile	GTC Val	AAA Lys 230	ACC Thr	TTG Leu	GTG Val	CAA Gln	GCT Ala 235	CAG Gln	AAT Asn	TCT Ser	AAA Lys	AGG Arg 240	720
35	CAC His	AAA Lys	GCC Ala	ATC Ile	CGT Arg 245	GTA Val	ATC Ile	ATA Ile	GCT Ala	GTG Val 250	GTG Val	CTT Leu	GTG Val	TTT	CTG Leu 255	GCT Ala	768
40	TGT Cys	CAG Gln	ATT Ile	CCT Pro 260	CAT His	AAC Asn	ATG Met	GTC Val	CTG Leu 265	CTT Leu	GTG Val	ACG Thr	GCT Ala	GCT Ala 270	AAT Asn	TTG Leu	816
45	GGT Gly	AAA Lys	ATG Met 275	AAC Asn	CGA Arg	TCC Ser	TGC Cys	CAG Gln 280	AGC Ser	GAA Glu	AAG Lys	CTA Leu	ATT Ile 285	Gly	TAT Tyr	ACG Thr	864
43	AAA Lys	ACT Thr 290	Val	ACA Thr	GAA Glu	GTC Val	CTG Leu 295	GCT Ala	TTC Phe	CTG Leu	CAC His	TGC Cys 300	TGC Cys	CTG Leu	AAC Asn	CCT Pro	912
50	GTG Val 305	Leu	TAC	GCT Ala	TTT Phe	ATT Ile 310	Gly	CAG Gln	AAG Lys	TTC Phe	AGA Arg 315	Asn	TAC Tyr	TTT	CTG Leu	AAG Lys 320	960
55	ATC Ile	TTG Leu	AAG Lys	GAC Asp	CTG Leu 325	Trp	TGT Cys	GTG Val	AGA Arg	AGG Arg 330	Lys	TAC	AAG Lys	TCC Ser	Ser 335	GTA	1008
60	TTC Phe	TCC	TGT Cys	GCC Ala 340	Gly	AGG Arg	TAC Tyr	TCA Ser	GAA Glu 345	Asn	ATT Ile	TCT Ser	CGG	Glr 350	Thr	AGT Ser	1056
	GAG Glu	ACC Thr	GCA Ala 355	Asp	AAC Asr	GAC Asp	TAA :	GCG Ala 360	Sex	TCC Ser	TTC Phe	ACT Thr	Met 365	:	\TAG#	AAG	1105

40

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CTGAGTCTCC CTAA

1119

כ	(2) INFORMATION FOR SEQ ID NO:10:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 365 amino acids
	(B) TYPE: amino acid
10	(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(ix) FEATURE:

15 (D) OTHER INFORMATION: Amino acid at position 193 may be His or Gln.

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

20 Met Phe Ser Thr Pro Val Lys Ile Ile Leu Cys Gln Ser Ile Leu His 1 5 10 15

Ile Thr Gln Leu Ile Leu Arg Cys Tyr Cys Ala Pro Cys Arg Arg Ser

20 . 25 30 30

Gly Ser Ser Pro Gly Tyr Leu Tyr Arg Ile Ala Tyr Ser Leu Ile Cys
35 40 45

Val Leu Gly Leu Leu Gly Asn Ile Leu Val Val Ile Thr Phe Ala Phe 30 50 55 60

Tyr Lys Lys Ala Arg Ser Met Thr Asp Val Tyr Leu Leu Asn Met Ala 65 70 75 80

35 Ile Ala Asp Ile Leu Phe Val Leu Thr Leu Pro Phe Trp Ala Val Ser 85 90 95

His Ala Thr Gly Ala Trp Val Phe Ser Asn Ala Thr Cys Lys Leu Leu 100 105 110

Lys Gly Ile Tyr Ala Ile Asn Phe Asn Cys Gly Met Leu Leu Leu Thr 115 120 125

Cys Ile Ser Met Asp Arg Tyr Ile Ala Ile Val Gln Ala Thr Lys Ser 45 130 135 140

50 Val Val Trp Gly Leu Ser Val Ile Ile Ser Ser Ser Thr Phe Val Phe 165 170 175

Asn Gln Lys Tyr Asn Thr Gln Gly Ser Asp Val Cys Glu Pro Lys Tyr 180 185 190

Gln Thr Val Ser Glu Pro Ile Arg Trp Lys Leu Leu Met Leu Gly Leu 195 200 205

Glu Leu Leu Phe Gly Phe Phe Ile Pro Leu Met Phe Met Ile Phe Cys
210 220

Tyr Thr Phe Ile Val Lys Thr Leu Val Gln Ala Gln Asn Ser Lys Arg 225 230 235 240

	.His	Lys	Ala	Ile	Arg 245	Val	Ile	Ile	Ala	Val 250	Val	Leu	Val	Phe	Leu 255	Ala.	
5	Cys	Gln	Ile	Pro 260	His	Asn	Met	Val	Leu 265	Leu	Val	Thr	Ala	Ala 270	Asn	Leu	
•	Gly	Lys	Met 275	Asn	Arg	Ser	Суз	Gln 280	Ser	Glu	Lys	Leu	Ile 285	Gly	Tyr	Thr	
10	Lys	Thr 290	Val	Thr	Glu	Val	Leu 295	Ala	Phe	Leu	His	Cys 300	Cys	Leu	Asn	Pro	
15	305					310					Arg 315					320	
13				•	325					330	Lys				335		
20				340					345		Ile			Gln 350	Thr	Ser	
			355					360		Ser	Phe	Thr	Met 365				
25	(2)						ID I										
		(i	(A) L	ENGT	H: 1	CTER	base	pai	rs							
30			ĺ	c) s	TRAN	DEDN	leic ESS: lin	sin	d gle								
		(ii) MC	LECU	LE T	YPE:	cDN	A									•
35		(ix	•	ATUR (A) N (B) I	IAME/	KEY:	CDS	.111	.6								
40		(xi	.) SI	QUE	NCE I	ESCF	RIPTI	ON:	SEQ	ID N	ю:11	.:					
	GAC	GAAC	CTG	CTT	GGGG	GG 1	CAGC	AAA:	T T	TTAF	DTAA	CAC	CAAAE	T AT Me	rG A7	C TAC	57
45				•											1		
	ACC Th:	r Arg	r TT(g Pho	c TT	A AAJ	A GG(s Gly	AGT Ser 10	Leu	AA(S ATC	G GCC	AAT AST	ı Tyı	C ACC	CTC C Let	G GCA	105
50	CC. Pr	o Gl	G GA	T GA	A TA' u Ty:	r GA' r Ası 2	p Val	CTY L Le	C AT	A GAJ e Gli	A GG? u Gly 30	y GI1	A CTO	G GA	G AG(u Se:	C GAT. r Asp 35	153
55	GA Gl	G GC. u Al	A GA a Gl	G CA u Gl	A TG n Cy 4	s As	C AAG	G TA' s Ty	T GA r As	C GCC p Ala	a Gl	G GC	A CT a Le	C TC u Se	A GCor Al.	C CAG a Gln 0	201
60	CT Le	G GT	G CC	o Se	A CT r Le	C TG u Cy	C TC	T GC r Al	a Va	G TT 1 Ph 0	T GT e Va	G AT	C GG e Gl	y va	C CT 1 Le 5	G GAC u Asp	249

	AAT Asn	CTC Leu	CTG Leu 70	Val	GTG Val	CTT Leu	ATC Ile	CTG Leu 75	Val	AAA Lys	TAT Tyr	AAA Lys	GGA Gly 80	Leu	AAA Lys	CGC Arg	2:	97
5	GTG Val	GAA Glu 85	TAA neA	ATC Ile	TAT Tyr	CTT	CTA Leu 90	AAC Asn	TTG Leu	GCA Ala	GTT Val	TCT Ser 95	AAC Asn	TTG Leu	TGT Cys	TTC Phe	34	45
10	TTG Leu 100	CTT Leu	ACC Thr	CTG Leu	CCC Pro	TTC Phe 105	TGG Trp	GCT Ala	CAT His	GCT Ala	GGG Gly 110	GGC Gly	GAT Asp	CCC Pro	ATG Met	TGT Cys 115	39	93
15	ьуs	iie	Leu	Ile	Gly 120	Leu	Tyr	Phe	Val	Gly 125	Leu	Tyr	AGT Ser	Glu	Thr 130	Phe	. 44	41
20	Pne	Asn	Суз	135	Leu	Thr	Val	Gln	Arg 140	Tyr	Leu	Val	TTT Phe	Leu 145	His	Lys	4.8	9
	GIĀ	Asn	150	Phe	Ser	Ala	Arg	Arg 155	Arg	Val	Pro	Cys	GGC Gly 160	Ile	Ile	Thr	53	37
25	Ser	165	Leu	Ala	Trp	Val	Thr 170	Ala	Ile	Leu	Ala	Thr 175	TTG Leu	Pro	Glu	Phe	58	15
30	180	Val	Тут	Lys	Pro	Gln 185	Met	Glu	Asp	Gln	Lys 190	Tyr	AAG Lys	Суѕ	Ala	Phe 195	63	13
35	ser	Arg	Thr	Pro	Phe 200	Leu	Pro	Ala	Asp	Glu 205	Thr	Phe	TGG Trp	Lys	His 210	Phe	. 68	1
40	Leu	Thr	Leu	Lys 215	Met	Asn	Ile	Ser	Val 220	Leu	Val	Leu	CCC Pro	Leu 225	Phe	Ile	72	9
45	Pne	Tnr	230	Leu	Tyr	Val	Gln	Met 235	Arg	Lys	Thr	Leu	AGG Arg 240	Phe	Arg	Glu	77	7
45	Gin	245	Tyr	Ser	Leu	Phe	Lys 250	Leu	Val	Phe	Ala	Val 255	ATG Met	Val	Val	Phe	82	5
50	260	Leu	Met	Trp	Ala	Pro 265	Tyr	Asn	Ile	Ala	Phe 270	Phe	CTG Leu	Ser	Thr	Phe 275	87	3
55	ràs	Glu	H1S	Phe	Ser 280	Leu	Ser	Asp	Cys	Lys 285	Ser	Ser	TAC Tyr	Asn	Leu 290	Asp	92:	1
60	Lys	ser	Val	His 295	Ile	Thr	Lys	Leu	11e 300	Ala	Thr	Thr	CAC His	Cys 305	Cys	Ile	96	9
	AAC Asn	Pro	CTC Leu 310	CTG Leu	TAT Tyr	GCG Ala	TTT Phe	CTT Leu 315	GAT Asp	GGG Gly	ACA Thr	TTT Phe	AGC Ser 320	AAA Lys	TAC Tyr	CTC Leu	101	7

	TGC CGC TGT TTC CAT CTG CGT AGT AAC ACC CCA CTT CAA CCC AGG GGG Cys Arg Cys Phe His Leu Arg Ser Asn Thr Pro Leu Gln Pro Arg Gly 325 330 335	1065
5	CAG TCT GCA CAA GGC ACA TCG AGG GAA GAA CCT GAC CAT TCC ACC GAA Gln Ser Ala Gln Gly Thr Ser Arg Glu Glu Pro Asp His Ser Thr Glu 340 345 350 355	1113
10	GTG TAAACTAGCA TCCACCAAAT GCAAGAAGAA TAAACATGGA TTTTCATCTT Val	1166
	TCTGCATTAT TTCATGTAAA TTTTCTACAC ATTTGTATAC AAAATCGGAT ACAGGAAGAA	1226
15	AAGGGAGAGG TGAGCTAACA TTTGCTAAGC ACTGAATTTG TCTCAGGCAC CGTGCAAGGC	1286
	TCTTTACAAA CGTGAGCTCC TTCGCCTCCT ACCACTTGTC CATAGTGTGG ATAGGACTAG	1346
20	TCTCATTTCT CTGAGAAGAA AACTAAGGCG CGGAAATTTG TCTAAGATCA CATAACTAGG	1406
20	AAGTGGCAGA ACTGATTCTC CAGCCCTGGT AGCATTTGCT CAGAGCCTAC GCTTGGTCCA	1466
	GAACATCAAA CTCCAAACCC TGGGGACAAA CGACATGAAA TAAATGTATT TTAAAACATA	1526
25	TAAAAAAAA AAAAAAAAA A	1547
30	(2) INFORMATION FOR SEQ ID NO:12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 356 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear	
35	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
40	Met Ile Tyr Thr Arg Phe Leu Lys Gly Ser Leu Lys Met Ala Asn Tyr 1 5 10 15	
	Thr Leu Ala Pro Glu Asp Glu Tyr Asp Val Leu Ile Glu Gly Glu Leu 20 25 30	
45	Glu Ser Asp Glu Ala Glu Gln Cys Asp Lys Tyr Asp Ala Gln Ala Leu 35 40 45	
50	Ser Ala Gln Leu Val Pro Ser Leu Cys Ser Ala Val Phe Val Ile Gly 50 55 60	
50	Val Leu Asp Asn Leu Leu Val Val Leu Ile Leu Val Lys Tyr Lys Gly 65 70 75 80	
55	Leu Lys Arg Val Glu Asn Ile Tyr Leu Leu Asn Leu Ala Val Ser Asn 85 90 95	
	Leu Cys Phe Leu Leu Thr Leu Pro Phe Trp Ala His Ala Gly Gly Asp 100 105 110	
60	Pro Met Cys Lys Ile Leu Ile Gly Leu Tyr Phe Val Gly Leu Tyr Ser 115 120 125	
	Glu Thr Phe Phe Asn Cys Leu Leu Thr Val Gln Arg Tyr Leu Val Phe 130 135 140	•

	Leu 145	His	Lys	Gly	Asn	Phe 150	Phe	Ser	Ala	Arg	Arg 155	Arg	Val	Pro	Суѕ	Gly 160	
5	Ile	Ile	Thr	Ser	Val 165	Leu	Ala	Trp	Val	Thr 170	Ala	Ile	Leu	Ala	Thr 175	Leu	
10	Pro	Glu	Phe	Val 180	Val	Tyr	Lys	Pro	Gln 185	Met	Glu	Asp	Gln	Lys 190	Tyr	Lys	
10	Сув	Ala	Phe 195	Ser	Arg	Thr	Pro	Phe 200	Leu	Pro	Ala	Asp	Glu 205	Thr	Phe	Trp	
15	Lys	His 210	Phe	Leu	Thr	Leu	Lys 215	Met	Asn	Ile	Ser	Val 220	Leu	Val	Leu	Pro	
	Leu 225	Phe	Ile	Phe	Thr	Phe 230	Leu	Tyr	Val	Glņ	Met 235	Arg	Lys	Thr	Leu	Arg 240	
20	Phe	Arg	Glu	Gln	Arg 245	Tyr	Ser	Leu	Phe	Lys 250	Leu	Val	Phe	Ala	Val 255	Met	
25	Val	Val	Phe	Leu 260	Leu	Met	Trp	Ala	Pro 265	Tyr	Asn	Ile	Ala	Phe 270	Phe	Leu	
23	Ser	Thr	Phe 275	Lys	Glu	His	Phe	Ser 280	Leu	Ser	Asp	Cys	Lys 285	Ser	Ser	Tyr	
30	Asn	Leu 290	Asp	Lys	Ser	Val	His 295	Ile	Thr	Lys	Leu	Ile 300	Ala	Thr	Thr	His	
	Cys 305	Cys ·	Ile	Asn	Pro	Leu 310	Leu	Tyr	Ala	Phe	Leu 315	Asp	Gly	Thr		Ser 320	
35	Lys	Tyr	Leu	Cys	Arg 325	Сув	Phe	His	Leu	Arg 330	Ser	Asn	Thr		Leu 335	Gln	
40	Pro	Arg	Gly	Gln 340	Ser	Ala	Gln	Gly	Thr 345	Ser	Arg	Glu		Pro 350	Asp	His	
40	Ser	Thr	Glu 355	Val													
45	(2)							10:13									
		(i)	(A	L) LE	ngth Pe :	: 35	5 am		acid	ls							
50			(1) TO	POLO	GY:	line		le								
		(ii)	MOL	ECUL	E TY	PE:	prot	ein									
55		(xi)	SEQ	UENC	E DE	SCRI	PTIC	N:S	EO I	D NO	:13:						
60												Asp	Thr	Thr	Thr	Glu 15	Phe
		Asp	Тут	Gly	Asp 20	Ala	Thr	Pro	Cys	Gln 25		Val	Asn	Glu		Ala	Phe
					-										30		

	Gly	Ala	Gln 35	Leu	Leu	Pro	Pro	Leu 40	Tyr	Ser	Leu	Val	Phe 45	Val	Ile	Gly
5	Leu	Val 50	Gly	Asn	Ile	Leu	Val 55	Val	Leu	Val	Leu	Val 60	Gln	Tyr	Lys	Arg
	Leu 65	Lys	Asn	Met	Thr	Ser 70	Ile	Tyr	Leu	Leu	Asn 75	Leu	Ala	Ile	Ser	Asp 80
10	Leu	Leu	Phe	Leu	Phe 85	Thr	Leu	Pro	Phe	Trp 90	Ile	Asp	Tyr	Lys	Leu 95	Lys
15	Asp	Asp	Trp	Val 100	Phe	Gly	Asp	Ala	Met 105	Cys	Lys	Ile	Leu	Ser 110	Gly	Phe
15	туг	Tyr	Thr 115	Gly	Leu	Tyr	Ser	Glu 120	Ile	Phe	Phe	Ile	11e 125	Leu	Leu	Thr
20	Ile	Asp 130	Arg	Tyr	Leu	Ala	Ile 135	Val	His	Ala	Val	Phe 140	Ala	Leu	Arg	Ala
	Arg 145	Thr	Val	Thr	Phe	Gly 150	Val	Ile	Thr	Ser	Ile 155	Ile	Ile	Trp	Ala	Leu 160
25	Ala	Ile	Leu	Ala	Ser 165	Met	Pro	Gly	Leu	Tyr 170	Phe	Ser	Lys	Thr	Gln 175	Trp
 30		Phe		180					185					190		•
30			195					200					205			
35		Leu 210					215					220				
	225					230					235					240
40		Phe			245					250			*		255	
45		Thr		260					265					270		
			275					280					285			Val
50		290	1				295					300				Val
	305	•				310	l				315					Val 320
55					325	,				330					335	
60	Ğlı	ı Arç	y Val	340		Thr	Ser	Pro	Ser 345		Gly	Glu	. His	350	Leu	Ser
	Ala	a G13	Phe 355													

(2) INFORMATION FOR SEQ ID NO:14:

5	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 374 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear														
	(ii)	MOLI	ECULI	E TY	PE: 1	prot	ein									
10																
	(xi)	SEQ	JENCI	E. DES	SCRI	PTIO	N: S	EQ II	D NO	:14:						
15	Met 1	Leu	Ser	Thr	Ser 5	Arg	Ser	Arg	Phe	Ile 10	Arg	Asn	Thr	Asn	Glu 15	Ser
20	Gly	Glu	Glu	Val 20	Thr	Thr	Phe	Phe	Asp 25	Tyr	Asp	тут	Gly	Ala 30	Pro	Cys
20	His	Lys	Phe 35	Asp	Val	Lys	Gln	Ile 40	Gly	Ala	Gln	Leu	Leu 45	Pro	Pro	Leu
25	Tyr	Ser 50	Leu	Val	Phe	Ile	Phe 55	Gly	Phe	Vạl	Gly	Asn 60	Met	Leu	Val	Val
	Leu 65	Ile	Leu	Ile	Asn	Cys 70	Lys	Lys	Leu	Lys	Cys 75	Leu	Thr	Asp	Ile	Tyr 80
30	Leu	Leu	Asn	Leu	Ala 85	Ile	Ser	Asp	Leu	Leu 90	Phe	Leu	Íle	Thr	Leu 95	Pro
35	Leu	Trp	Ala	His 100	Ser	Ala	Ala	Asn	Glu 105	Trp	Val	Phe	Gly	Asn 110	Ala	Met
	Суѕ	Lys	Leu 115	Phe	Thr	Gly	Leu	Tyr 120	His	Ile	Gly	Tyr	Phe 125	Gly	Gly	Ile
40	Phe	Phe 130	Ile	Ile	Leu	Leu	Thr 135	Ile	Asp	Arg	Tyr	Leu 140	Ala	Ile	Val	His
	Ala 145	Val	Phe	Ala	Leu	Lys 150	Ala	Arg	Thr	Val	Thr 155	Phe	Gly	Val	Val	Thr 160
45	Ser	Val	Ile	Thr	Trp 165	Leu	Val	Ala	Val	Phe 170	Ala	Ser	Val	Pro	Gly 175	Ile
50	Ile	Phe	Thr	Lys 180	Суз	Gln	Lys	Glu	Asp 185	Ser	Val	Tyr	Val	Cys 190	Gly	Pro
50	Tyr	Phe	Pro 195	Arg	Gly	Trp	Asn	Asn 200	Phe	His	Thr	Ile	Met 205	Arg	Asn	Ile
55	Leu	Gly 210	Leu	Val	Leu	Pro	Leu 215	Leu	Ile	Met	Val	Ile 220	Cys	Tyr	Ser	Gly
•	Ile 225	Leu	Lys	Thr	Leu	Leu 230	Arg	Cys	Arg	Asn	Glu 235	Lys	Lys	Arg	His	Arg 240
60	Ala	Val	Arg	Val	Ile 245	Phe	Thr	Ile	Met	Ile 250	Val	Tyr	Phe	Leu	Phe 255	Trp
	Thr	Pro	Tyr	Asn 260	Ile	Val	Ile	Leu	Leu 265	Asn	Thr	Phe	Gln	Glu 270	Phe	Phe

	Gly	Leu	Ser 275	Asn	Cys	Glu		Thr 280	Ser	Gln	Leu	Asp	Gln 285	Ala	Thr	Gln
5	Val	Thr 290	Glu	Thr	Leu		Met 2 9 5	Thr	His	Cys	Cys	Ile 300	Asn	Pro	Ile	Ile
10	туг 305	Ala	Phe	Val	Gly	Glu 310	Lys	Phe	Arg	Ser	Leu 315	Phe	His	Ile	Ala	Leu 320
10	Gly	Cys	Arg	Ile	Ala 325	Pro	Leu	Gln		Pro 330	Val	Cys	Gly	Gly	Pro 335	Gly
15	Val	Arg	Pro	Gly 340	Lys	Asn	Val	Lýs	Val 345	Thr	Thr	Gln	Gly	Le u 350	Leu	Asp
	Gly	Arg	Gly 355	Lys	Gly	Lys		Ile 360	Gly	Arg	Ala	Pro	Glu 365	Ala	Ser	Leu
20	Gln	Asp 370	Lys	Glu	Gly	Ala										
(2) INFORMATION FOR SEQ ID NO:15:																
25	(i)	(B)	LEN	WGTH:	: 35! mino	ami aci	no a	cids	3							
30	(C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein															
35	(xi)	SEQ	JENCI	E DE	SCRI:	PTIO	N: SI	EQ II	O NO:	:15:						
. •											Phe	Gly	Thr	Thr	Ser 15	Tyr
35 40	Met 1		Thr	Ser	Leu 5	Asp	Thr	Val	Glu	Thr 10					15	
. •	Met 1 Tyr Met	Thr Asp	Thr Asp Gln 35	Ser Val 20	Leu 5 Gly Val	Asp Leu Pro	Thr Leu Pro	Val Cys Leu 40	Glu Glu 25 Tyr	Thr 10 Lys Ser	Ala Leu	Asp Val	Thr Phe 45	Arg 30 Thr	Ala Val	Leu
40	Met 1 Tyr Met	Asp Ala Leu 50	Asp Gln 35 Gly	Val 20 Phe	Leu 5 Gly Val	Asp Leu Pro Val	Thr Leu Pro Val	Val Cys Leu 40 Val	Glu 25 Tyr Met	Thr 10 Lys Ser	Ala Leu Leu	Asp Val Ile 60	Thr Phe 45	Arg 30 Thr	Ala Val Arg	Leu Gly Arg
40	Met 1 Tyr Met Let 65	Thr Asp Ala Leu 50	Thr Asp Gln 35 Gly	Val 20 Phe Asn	Leu 5 Gly Val Val	Asp Leu Pro Val Asn 70	Thr Leu Pro Val 55	Val Cys Leu 40 Val	Glu 25 Tyr Met Leu	Thr 10 Lys Ser Ile	Ala Leu Leu Asn 75	Asp Val Ile 60 Leu	Thr Phe 45 Lys	Arg 30 Thr Tyr	Ala Val Arg	Leu Gly Arg Asp
40 45 50	Met 1 Tyr Met Let 65	Asp Ala Leu 50	Thr Asp Gln 35 Gly	Val 20 Phe Asn	Leu 5 Gly Val Val	Asp Leu Pro Val Asn 70	Thr Leu Pro Val 55	Val Cys Leu 40 Val	Glu 25 Tyr Met Leu	Thr 10 Lys Ser Ile	Ala Leu Leu Asn 75	Asp Val Ile 60 Leu	Thr Phe 45 Lys	Arg 30 Thr Tyr	Ala Val Arg	Leu Gly Arg Asp
4 0	Met Tyr Met Let 65	Thr Asp Ala Leu 50	Asp Gln 35 Gly Ile	Val 20. Phe Asn Met	Leu 5 Gly Val Val Val 85	Asp Leu Pro Val Asn 70	Thr Leu Pro Val 55 Ile	Val Cys Leu 40 Val Tyr	Glu 25 Tyr Met Leu	Thr 10 Lys Ser Ile Leu Trp 90	Leu Leu Asn 75	Asp Val Ile 60 Leu His	Thr Phe 45 Lys Ala	Arg 30 Thr Tyr Ile	Ala Val Arg Ser Arg 95	Leu Gly Arg Asp 80
40 45 50	Met Tyr Met Let 65 Let	Thr Asp Ala Leu 50 Arg Leu Arg	Asp Gln 35 Gly Ile Phe Tr; Thr	Val 20 Phe Asm Met	Leu 5 Gly Val Val Val 85 Phe	Asp Leu Pro Val Asn 70 Thr	Thr Leu Pro Val 55 Ile Leu His	Val Cys Leu 40 Val Tyr Pro Gly Glu 120	Glu 25 Tyr Met Leu Phe 105	Thr 10 Lys Ser Ile Leu Trp 90 Cys	Ala Leu Leu Asn 75 Ile	Val Ile 60 Leu His	Thr Phe 45 Lys Ala Tyr	Arg 30 Thr Tyr Ile Val	Ala Val Arg Ser Arg 95 Gly	Leu Gly Arg Asp 80

		145	THE	vai	Thr	Pne	150	۸ġŢ	116	Thr	Ser	11e 155	Val	Thr	Trp	Gly	160
5		Ala	Val	Leu	Ala	Ala 165	Leu	Pro	Glu	Phe	Ile 170	Phe	Tyr	Glu	Thr	Glu 175	Glu
		Leu	Phe	Glu	Glu 180	Thr	Leu	Сув	Ser	Ala 185	Leu	Tyr	Pro	Glu	Asp 190	Thr	Val
10		Тут	Ser	Trp 195	Arg	His	Phe	His	Thr 200	Leu	Arg	Met	Thr	Ile 205	Phe	Cys	Leu
15		Val	Leu 210	Pro	Leu	Leu	Val	Met 215	Ala	Ile	Cys	Tyr	Thr 220	Gly	Ile	Ile	Lys
		Thr 225	Leu	Leu	Arg	Cys	Pro 230	Ser	Lys	Lys	Lys	Tyr 235	Lys	Ala	Ile	Arg	Leu 240
20		Ile	Phe	Val	Ile	Met 245	Ala	Val	Phe	Phe	Ile 250	Phe	Trp	Thr	Pro	Tyr 255	Asn
		Val	Ala	Ile	Leu 260	Leu	Ser	Ser	Tyr	Gln 265	Ser	Ile	Leu	Phe	Gly 270	Asn	Asp
25		Cys	Glu	Arg 275	Ser	Lys	His	Leu	Asp 280	Leu	Val	Met	Leu	Val 285	Thr	Glu	Val
30		Ile	Ala 290	Tyr	Ser	His	Cys	Cys 295	Met	Asn	Pro	Va1	Ile 300	Tyr	Ala	Phe	Val
		Gly 305	Glu	Arg	Phe	Arg	Lys 310	Tyr	Leu	Arg	His	Phe 315	Phe	His	Arg	His	Leu 320
35		Leu	Met	His	Leu	Gly 325	Arg	Tyr	lle	Pro	Phe 330	Leu	Pro	Ser	Glu	Lys 335	Leu
		Glu	Arg	Thr	Ser 340	Ser	Val	Ser	Pro	Ser 345	Thr	Ala	Glu 、	Pro	Glu 350	Leu	Ser
40		Ile	Val	Phe 355					-	•							
	(2)	INFOR	RMATI	ON E	FOR S	SEQ 1	D NO	0:16:	;								
45		(i)	(A)	LEN	NGTH: PE: 8	ARACT : 360 amino	ami aci	ino a	cids	3							
50	•					3Y:]											
		(ii)	MOLI	3CULI	TYI	PE: p	rote	in				•					
55	,	(xi)	SEQ	JENCI	E DES	SCRI	PTIO	N: SI	ii Q	NO:	:16:		•				
<i>6</i> 0		Met 1	Asn	Pro	Thr	Asp 5	Ile	Ala	Asp	Thr	Thr 10	Leu	Asp	Glu	Ser	Ile 15	Туг
60		Ser	Asn	Tyr	Tyr 20	Leu	Tyr	Glu	Ser	Ile 25	Pro	Lys	Pro	Cys	Thr 30	Lys	Glu

	Gly		Lys 35	Ala	Phe	Gly	Glu	Leu 40	Phe	Leu	Pro	Pro	Leu 45	Tyr	Ser	Leu
5	Val	Phe 50	Val	Phe	Gly		Leu 55	Gly	Asn	Ser	Val	Val 60	Val	Leu	Val	Leu
	Phe 65	Lys	Tyr	Lys	Arg	Leu 70	Arg	Ser	Met	Thr	Asp 75	Val	Тут	Leu	Leu	Asn 80
10	Leu .	Ala	Ile		Asp 85	Leu	Leu	Phe	Val	Phe 90	Ser	Leu	Pro	Phe	Trp 95	Gly
15	Tyr			100					105					110		
	Ile		115	٠				120					125			*
20		130					135					140				
	Ser 145					150					155					160
25	Thr				165					170					175	
30	Thr			180					185		•			190		
			195				-	200					205	Ile		
35		210					215					220	•	Tyr		Ala
	225					230					235					240
40					245					250				Phe	255	
45		-		260					265					270		
			275					280					285			Ala
50		290					295					300				Tyr
	305					310	,	•			315					Lys 320
55					325	i				330					335	Gln Met
60				340)				345		-3-		J 21.	350)	-3
	Asp	HIS	355	Lev 5	. HIS	. WRI	, WT	360								

(2) INFORMATION FOR SEQ ID NO:17:

5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
	GTAATGATCA GTCAACGGGG GAC	. 23
15	(2) INFORMATION FOR SEQ ID NO:18:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid	
20	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	,
	CCAGCAAGCT TGCAACCTTA ACCA	24
	(2) INFORMATION FOR SEQ ID NO:19:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 9 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single	
35	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: peptide	
40		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	
	Asp Tyr Lys Asp Asp Asp Lys Leu 1 5	
45		

WHAT IS CLAIMED IS:

- A substantially pure or isolated polypeptide comprising a segment exhibiting sequence homology to a
 corresponding portion of a mature protein selected from the group consisting of:
 - i) TECK;
 - ii) MIP-3α;
 - iii) MIP-3β;
- 10 iv) DC CR; and
 - v) M/DC CR;

wherein said homology is at least about 70% identity and said portion is at least about 25 amino acid residues.

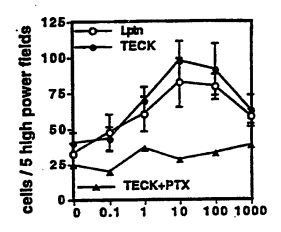
- The polypeptide of claim 1, further comprising a second segment exhibiting:
 - a) at least about 90% identity over at least 9 amino acid residues; or
- b) at least about 80% identity over at least 17
 amino acid residues.
 - 3. The polypeptide of either claim 1 or 2, selected from the group consisting of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 8, SEQ ID NO: 10 and SEQ ID NO: 12.
 - 4. An isolated or recombinant nucleic acid which encodes the polypeptides of any of claims 1-3.
- 30 5. The nucleic acid of claim 4, selected from the group consisting of SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7, SEQ ID NO: 9, and SEQ ID NO: 11.
- 6. A nucleic acid which hybridizes under stringent wash conditions of 55° C and less than 150 mM salt to a nucleic acid of claim 4 or 5.
 - 7. The nucleic acid of claim 6, which further exhibits at least about 85% identity over a stretch of at

least about 30 nucleotides to a nucleic acid of claim 4 or 5.

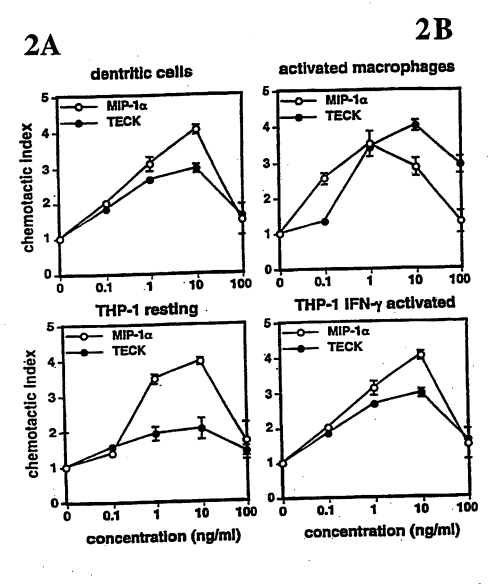
- 8. A vector comprising a nucleic acid of any of claims 4-7.
 - 9. A host cell comprising a nucleic acid or vector of any of claims 4-8.
- 10 10. A method for making a polypeptide comprising culturing a host cell of claim 9 under conditions in which the nucleic acid or vector is expressed.
- 11. A binding compound comprising an antibody or antigen binding fragment therefrom which binds to a polypeptide of any of Claims 1-3.

1/2

1 A



concentration (ng/ml)



2C

2D

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